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8 Statement of Inventorship
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Description	-
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11 I/We request the grant of a patent on the basis of this application.

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18 December 1998

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IMMUNOSUPPRESSION

1. FIELD OF THE INVENTION

5 This invention relates to immunosuppression and, more particularly, to immunosuppression in the context of xenotransplantation.

2. BACKGROUND TO THE INVENTION

10 Despite the established success of allogeneic organ transplantation, the increasing disparity between the supply and demand of organs must be overcome. Increasing the supply of allogeneic organs does not offer a satisfactory solution because even if all usable organs were transplanted this would still not meet the existing demand (1,2). This 15 has led to a resurgence of interest in xenotransplantation (the transplantation of organs between animals of different species) as a viable and attractive alternative.

20 Xenotransplantation research has recently focused on the pig as a suitable animal donor in terms of size, physiological compatibility and breeding characteristics (3,4). Until recently however, discordant xenotransplantation has been limited by the inevitable occurrence of humorally-mediated hyperacute rejection (HAR) which rapidly triggers 25 organ rejection upon revascularisation. HAR is the fate of most organs transplanted between discordant species. Recently, significant advances have been made in understanding the immunological basis of HAR, and many approaches have been employed to overcome it. Of significance, a variety of transgenic strategies are currently 30 being employed including the expression of regulators of complement activity on porcine endothelial cells (5). It is foreseeable that short-term xenograft survival will soon be achieved (6). The recent advances in overcoming HAR have highlighted subsequent immunological barriers which must be surmounted to enable long-term xenograft survival. Both humoral and cellular arms of the immune response appear to play a role in the downstream events of immunological rejection. Clearly the most important of which 35 is the existence of a formidable T cell mediated rejection response (7-11) previously

obscured by the dominant role of HAR. *In vitro*, human T cells have been demonstrated to play a central role in the recognition of xenogeneic cells (7,8,12) following sensitisation via the direct and indirect T cell activation pathways, which have been well documented for allore cognition and allograft rejection (13). Knowledge of the cellular mechanisms underlying allorejection has provided an important basis for the investigation of the T cell mediated xenoresponse.

At present, the major therapies to prevent cell mediated rejection of organ transplants rely on systemic immunosuppressive drugs or monoclonal antibody (Mab) therapy directed against targets such as CD3, CD4, CD25, (14). Following reports that strong T cell xenoresponses can be generated *in vitro* (7,8,12), control of xenograft rejection may require levels of immunosuppression much greater than the current standard doses. Such a strategy would not be desired in a xenograft context. Drugs must be taken for life, depress the entire immune system and result in an increased risk of infection and susceptibility to cancer (14). For the applicability of xenotransplantation to the clinic, targeting graft-specific strategies for tolerance induction/immunosuppression would clearly be highly advantageous. Whilst this has been difficult to achieve in an allotransplant context, xenotransplantation offers greater potential - with differences between species providing the option for the generation of reagents that are truly graft specific. In addition, there is the opportunity for the manipulation of both the porcine donor organ, and the human recipient's immune system, prior to transplantation (1).

3. DETAILED BACKGROUND

3.1 T cell activation and proliferation

Optimal proliferation of T cells, although initiated via ligation of the antigen specific CD3/TCR complex (Signal 1) requires additional costimulatory signals (Signal 2) (15,16,17) which are usually supplied by the antigen presenting cell (APC). Whilst antigenic stimulation of T cells in the presence of signal 2 induces T cell activation and proliferation (18), exposure of T cells to MHC-antigen complexes in their absence leads to aborted T cell proliferation and the development of clonal anergy (19,20).

Manipulation of APC by aldehyde fixation (20,21) or heat treatment (19) has been demonstrated to abrogate the ability of such cells to activate alloreactive T cells, without altering levels of MHC-II surface expression. Thus T cell receptor occupancy alone is insufficient to fully activate the T cell (17). Anergic T cells are best characterised by their

5 lack of IL-2 production and their continued inability to produce IL-2 on subsequent exposure to antigen (22). Thus, confirming the two signal model of activation as predicted by Lafferty *et al* (23). For T cells to respond to a given antigenic stimulus, multiple activation signals are required from the APC (23).

● 10 The *in vivo* induction of T cell anergy in the absence of a secondary signal was first demonstrated by Jenkins and Schwartz in 1986 (24) using chemically fixed APC to present specific peptide to CD4 T helper clones. A multitude of *in vitro* and *in vivo* data has since been produced supporting the hypothesis that signal 1 in isolation fails to activate T cells (22), and that costimulatory signalling results from contact with other

15 cells rather than via soluble factors. Fibroblasts transfected with human Class II MHC molecules, but not expressing the appropriate CS signals (lacking signal 2) can efficiently present antigen to class II restricted CD4 T cell clones, but these fail to cause antigen specific T cell proliferation, rendering cells anergic. The context in which T cells first encounter antigen therefore has an important bearing on subsequent immune

● 20 responsiveness.

Thus, costimulatory molecules are essential for T cell activation and multiplication and result from interactions between receptors on T cells and their ligands expressed on the APC. The costimulatory signal itself, however, is neither antigen specific nor MHC restricted (25). In recent years the molecular interactions involved in mediating costimulation have been well defined. The two key pathways involve (i) B7-1, B7-2 (members of the B7 family) and (ii) CD40, which are expressed on the APC, and their counter-receptors CD28 and CD40 ligand (CD40L) respectively expressed on T cells. A large body of evidence, both *in vivo* and *in vitro*, clearly defines the crucial roles played

25 30 by B7-1, B7-2 and CD40 in providing T cell costimulation (26-36). Furthermore, the

simultaneous blockade of signalling via CD28-B7 and CD40-CD40L in an allotransplant context prevented the onset of allograft rejection (37,38). *In vivo*, targeting the B7/CD28 interaction has been shown to prevent T cell sensitisation to graft antigen, thereby prolonging graft survival (38,39).

5

T cells can be sensitised against xenoantigens via one of two pathways - the direct and indirect pathways, which are analogous to the well documented T cell activation pathways against alloantigens (Figure 1). Direct recognition requires that the recipient T cells recognise intact xeno MHC-molecules complexed with peptide on donor stimulator 10 cells. In contrast, indirect recognition requires that recipient APC process the xenoantigen prior to presentation to recipient T cells in the context of recipient MHC II. Self MHC II restricted T cells with specificity for the xenoantigen will recognise the peptide and respond. Whilst the majority of data reported is of indirect xenorecognition responses, cell mediated rejection via the direct route has also been documented 15 (7,8,9,11,12,40,41,42). Vigorous human T cell proliferative responses directed against porcine tissues *in vitro* have been documented from studies both in this laboratory and others.

3.2 Costimulatory molecules

20 The crucial role played by costimulatory molecules in determining the result of TCR-CD3 receptor engagement with MHC and peptides has been demonstrated extensively both *in vivo* and *in vitro*. Anti-costimulatory molecule strategies aimed at either the receptors or their ligands are being used as therapeutic strategies for altering the immune response. Such approaches have been tested in model transplant systems to alter cell 25 mediated responses thereby preventing graft rejection (14,37,38,43-47).

B7-1 (B7/BB1, CD80) and B7-2 (CD86) both belong to the Immunoglobulin superfamily and are heavily glycosylated transmembrane proteins (25). B7-1, a B cell activation molecule was first identified in 1981 (27), followed by B7-2 in 1993 (49). Both human 30 B7-1 and B7-2, and the murine homologues have now been cloned and functionally

characterised (25). B7-1 and B7-2 are constitutively expressed on splenic and blood dendritic cells and are induced on B cells and monocytes upon activation (34,50.). B7-1 and 2 are highly homologous and are the natural ligands for the T cell antigen CD28 (50). Cytotoxic T lymphocyte antigen-4 (CTLA-4), a cell surface glycoprotein has been 5 identified as a second receptor for the B7 family of molecules (51) and is homologous to CD28 with 31% sequence identity. Both B7 isoforms bind to CTLA-4 with higher affinity than to CD28 (30,50,52). Whilst CD28-B7 receptor engagement results in an APC-derived costimulatory signal involved in antigen specific IL-2 production both *in vivo* and *in vitro* (53,54), CTLA4 appears to function as a negative regulator of T cell activation (55, 56, 57). Cross-linking by anti-CTLA4 antibodies has been demonstrated to antagonise CD28 ligation (58) and, in addition, CTLA4 knock-out mice die due to uncontrolled lymphocyte proliferation within the first few weeks of life (59). Thus, CTLA4 ligation is thought to be crucial for the maintenance and regulation of immune responses. The underlying mechanisms have not, however, been clearly defined.

15 Among costimulatory molecules, the B7 family appears to be unique, since ligation by CD28 of either B7-1 or B7-2 is both necessary and sufficient to prevent the induction of anergy (34). The CD28-B7 interaction is thought to deliver crucial signals to sustain proliferation of activated T cells. These observations are supported by *in vitro* data 20 showing that whilst cells deficient in B7 fail to stimulate a primary MLR, transfectants expressing high levels of B7 gained the capacity to stimulate the production of IL-2 by alloreactive T cells and to co-stimulate a polyclonal population of purified T cells cultured with immobilised anti-CD3 Mab (31). Artificial APC generated by stably transfecting NIH-3T3 cells with HLA-DR7, B7 or both, clearly demonstrated that following 25 presentation of tetanus toxoid (TT) optimal T cell proliferation and IL-2 production resulted only when both molecules were present. In the absence of B7, clonal anergy resulted (58).

30 Porcine B7-2 (PoB7-2) has been cloned from aortic endothelial cells (60). Following transient transfection of porcine B7-2, human umbilical vein endothelial cells strongly

costimulated IL-2 production by human T cells. This costimulation of human T cells by poB7-2 was shown to be as effective as costimulatory signals provided by human B7-1 or B7-2 and could be specifically blocked by huCTLA4Ig. Thus poB7-2 strongly contributes to the immunogenicity of porcine endothelium (60).

5

Although B7-1 and B7-2 mediated interactions appear to be central to the development of T cell specific immunity, additional costimulatory pathways of importance exist. The most crucial of which involves the CD40 and CD40 ligand (CD40L) interaction (34).

10 CD40 is a 50kDa surface glycoprotein belonging to the TNF-receptor superfamily. CD40 is expressed on various APC including among others, monocytes, dendritic cells and activated macrophages. Other cell types including endothelium also express CD40 (34). Its counter-receptor CD40L (CD154, gp39, TRAP) is a 33 kDa type II integral membrane protein (34,36) transiently expressed on activated CD4 T cells. The CD40-CD40L 15 interaction has been demonstrated to play an important role in both the humoral and cellular arms of the immune response with a dominant role in B cell activation. Whilst cross linking of CD40 on B cells is essential for B cell growth and isotype switching, it also results in the upregulation of B7 expression (50). Levels of B7 expression (and thus 20 APC capacity) of monocytes and dendritic cells are clearly unregulated following CD40 signalling (34). Data from CD40 knock-out mice demonstrated that CD40L signalling following ligation by CD40 plays an important role in T cell activation (61). Transfection of the murine P815 mastocytoma cells with CD40 (or B7-1) enabled previously non-stimulatory P815 cells to mediate the costimulation necessary for polyclonal T cell activation and the generation of cytokines (34). CD40-CD40L interactions have also been 25 demonstrated to play a critical role in allograft rejection (62,63).

Resting B cells do not normally express B7-1/B7-2 at high levels until they are activated (50). Activation of B cells following simultaneous engagement of MHC-peptide/TCR and CD40-CD40L leads to the upregulation of B7 family members on B cells, thereby 30 enhancing the stimulation and subsequent activation of T cells (34,36). Thus, the

CD40-CD40L interaction influences costimulatory activity by inducing expression of the B7 family of molecules and perhaps other costimulatory molecules, thereby playing a key role in T cell activation. The clear synergistic effects of CD40 and B7 indicate the importance of both costimulatory pathways for the initiation and amplification of T cell dependent immune responses (38). CD40-CD40L interactions have also been shown to play a crucial role in the generation of cytotoxic T lymphocyte (CTL) responses by modifying the functional status of a dendritic cell (64,65,66)

Extensive studies have demonstrated the importance of blocking B7-CD28 and/or CD40-CD40L interactions in the context of both allo and xenotransplantation. Data strongly supporting this includes the use of CTLA4Ig to block signalling via CD28-B7 resulting in enhanced graft survival and the prevention of chronic rejection in a rat cardiac allograft model (44,45) and a murine aortic allograft model (43). In these models, administration of CTLA4Ig caused partial (44) or complete (46) tolerance to graft antigen by inducing T cell anergy. Treatment of allo pancreatic islet transplants with anti-B7-2 and B7-1 antibody has also been demonstrated to inhibit transplant rejection (14). Similar results were obtained in models inhibiting CD40 signalling in a mouse cardiac allotransplant models (37,47,62). Two studies detailing the simultaneous blockade of signalling via CD28-B7 and CD40-CD40L prevented the onset of allorejection. Concurrent prolonged inhibition of both pathways completely abrogated the onset of chronic rejection in a mouse allo model (37) and in a skin and heart allo model (38).

In the realm of xenotransplantation, Lenschow and colleagues have, demonstrated long-term donor specific tolerance of human islets transplanted into mice with concomitant treatment with CTLA4Ig (46). Graft specific tolerance was demonstrated to be a direct consequence of inhibiting recognition via B7 expressing APC. In addition, Tran *et al* (67) demonstrated short term suppression with CTLA4-Fc treatment. There is limited data available on the simultaneous blockade of both pathways in the xenotransplantation context, with the prolonged survival of rat and porcine skin transplanted into murine recipients (63).

In vitro and *in vivo* data have clearly demonstrated that targeting the interactions mediated by either the CD28-B7, CD40-CD40L, or both pathways has prevented the sensitisation of T cells to alloantigen and xenoantigen from engrafted tissue thereby 5 prolonging graft survival (6).

3.3 Peptide immunisation strategy

Previous *in vivo* studies using synthetic peptides conjugated to carrier molecules as immunogens have demonstrated the ability to generate the production of biologically 10 active antibodies (68). There is now an extensive literature detailing peptide immunisation strategies which demonstrate enhancement of antibody production by carrier presentation(68-72). Thus, appropriate T cell epitopes can be used to prime T cells for subsequent help to B cells. Recent data has been published reporting the production of IgG by self-reactive B cells following immunisation with a self reacting antigen 15 covalently coupled to a carrier molecule (70). Thereby demonstrating that B cell tolerance to self protein can be overcome.

As mentioned above, in order to be recognised by T cells, antigen (self or foreign) must be processed and presented by APC. B cells can act as highly potent APC following 20 endocytosis of antigen via IgG receptors . In the presence of a full complement of activation signals (TCR engagement plus costimulation) T cell activation will occur resulting in the subsequent generation of antibody.

Peptides from self proteins are processed and presented to T cells in the same manner as 25 foreign proteins, but because of T cell tolerance, presentation of self peptides does not normally result in T cell activation (70). The absence of T cell recognition may therefore explain, in part, why potentially reactive B cells fail to respond.

The ability to overcome B cell non-responsiveness to self peptides has recently been 30 demonstrated by Dalum *et al* (69). An autoantibody response was generated by the

provision of additional T cell help in the form of a strong foreign carrier T cell epitope. Further studies have demonstrated that synthetic peptides conjugated to T cell carrier molecules are capable of overcoming B cell non-responsiveness if significant numbers of self-reactive B cells are present in the host (69,70). Insertion of a single foreign T cell 5 epitope into the sequence of Ubiquitin, elicited strong autoantibody production directed against the native molecule (69). In an elegant study by Sad, using GnRH as a self protein chemically linked to diphtheria toxoid (DT) as the synthetic T cell epitope, autoantibodies were produced with specificity for native GnRH (71,72). Following the initial vaccination, the continued presence of the native GnRH *in vivo* maintained the 10 production of Ab. Continued antibody production caused sterility in the immunised mice due to the sustained anti-GnRH antibody response maintained by the continued presence of the native molecule against which the specific B cells were producing antibody. The DT carrier provoked a helper T cell response to assist GnRH specific B cells and break B cell tolerance.

15

4. STATEMENTS OF INVENTION

The present invention involves the use of a foreign T cell carrier to exert significant influences on subsequent responses to molecules conjugated to the carrier. By such 20 means autoantibody responses may be directed against costimulatory molecules in a xenotransplantation context.

According to the present invention there is provided a method of improving the tolerance of an animal, including a human being, to a xenograft, the animal having T cell mediated 25 immunity, the method comprising causing the animal to raise an antibody against a xenomecuae involved in the general of a rejection response in the animal, said antibody being raised by immunising the animal with a chimeric peptide comprising a T cell epitope against which the animal has immunity and a B cell epitope of said xenomecuae.

30 Accordingly, xenograft specific tolerance is induced in transplant recipients by targeting the direct T cell mediated response by the use of chimeric peptide constructs to stimulate

the generation of specific anti-graft tolerance-promoting antibodies by the recipient prior to transplantation. By way of example, the chimeric peptides comprise a T cell epitope conjugated to sequences of porcine costimulatory molecules, B7-1, B7-2 and CD40. The presence of the engrafted tissue will then serve to maintain and perpetuate the production 5 of antibody by the recipient's B cells.

The present invention also provide a chimeric peptide comprising a T cell epitope and a B cell epitope, said T cell being that of an animal, including a human being of a first species and said B cell being of an animal of a second species, said first and second species such 10 that xeno transplantsations suitable from an animal of said second species to an animal of said first species.

In addition, the present invention provides the use of a chimeric peptide improving the tolerance of an animal, including a human being, to a xenograph, the chimeric peptide 15 being as defined above.

The potential benefits of the use of a chimeric peptide of the invention are that it avoids the need for injection of blocking antibodies or fusion proteins. Furthermore, the induction of a recipient antibody response circumvents the problems most commonly 20 associated with administration of xenogeneic antibodies or fusions proteins, namely the immune response against the administered reagent.

5. SPECIFIC EMBODIMENTS

5.1 Cloning porcine costimulatory molecules

25 5.1.1 Cloning porcine B7-2

RNA was extracted from primary and transformed porcine cells using a standard protocol. mRNA was then reverse transcribed and porcine B7-2 (poB7-2) amplified from the cDNA by 35 cycles of PCR at 56° C with 1.5mM magnesium. The 5' and 3' primers GCATGGATCCATGGGACTGAGTAACATTCTTTG and 30 GCATGTCGACTTAAAAATCTGTAGTACTGTTGTC respectively were designed on

the basis of the published poB7-2 sequence (60) to overlay the start and stop codons (Figure 2). A 956 base pair fragment was generated and subcloned into the BamH1 & Sal1 restriction sites of pbluescript. The nucleotide sequence was determined using standard m13 forward and reverse primers. The sequence of a single clone, CD86(i) is 5 illustrated in Figure 3, with comparison to the published sequences from porcine (Figure 4), human and murine B7-2 (Figure 5). One base pair difference is detected between our clone, CD86(i), and the published sequence at the 3' prime end. This, however, is unlikely to be an important difference with respect to either poB7-2 expression or binding to its ligand. The predicted amino acid sequence of CD86(i), compared to that of porcine, 10 human and mouse B7-2 is shown in Figure 6.

5.1.2 Cloning porcine B7-1 and CD40

RNA extracted from phytohaemagglutinin (PHA) or poke-weed mitogen (PMW) stimulated porcine PBMC and transformed porcine endothelial cells is being used to 15 amplify cDNA encoding the costimulatory molecules B7-1 and CD40. B7-1 Primers were designed on the basis of conserved areas following comparison of murine and human (29,49) sequences. External (lying outside the coding region) AGACCGTCTCCTTTAG(3'i), TTGGATCCTCCATGTTATCCC (3'ii) and 20 AGCATCTGAAGC (5') and internal (within the coding region) ATGGATCCTCCATTTCCAACC (3') and TTGTCGACATCTACTGGC (5') primers have been designed as depicted in Figure 7. The generation of two 3' primers is due to significant differences between the human and murine sequences in the terminal coding regions. Resulting PCR fragments will be subcloned as described above using the restriction sites BamHI and SalI contained within the promoter sequence. Constructs will 25 then be sent for sequence confirmation.

CD40 primers were designed in a similar manner following sequence alignment of published CD40 sequences from human, mice and cattle (73,74,75) as illustrated in Figures 8A & B. The 5' and 3' primer sequences are 30 GGATCCTCACTGTCTCCTGCAGTGCGACTCTCCTTTGCCGTCCG

TCCTCC and GAATTCATGGTCTGTTGCCTCTGCAGTG respectively containing the BamHI and EcoRI restriction sites.

5.2 Generation of porcine costimulatory molecule expressing cell transfectants

5 The poB7-2 molecule (CD869(i)) has been subcloned into the eukaryotic expression vector pci.neo carrying the neomycin drug-selectable marker. This is being used to transfect M1 and M1.DR1 transformed murine cell lines using a standard calcium phosphate precipitation method. G418 resistant pci.neo expressing cells will be selected using dynabead purification and highly expressing clones will be selected by limiting 10 dilution.

Stable poB7-2 M1 and P815 transfectants have been generated by this approach using the poB7-2 DNA construct supplied to us by Maher *et al* (Figure 9). transient transfections of M1 and P815 cells have been generated using our CD86(i) construct (Figure 10).

3 particular assays will be undertaken using the CD86(i) transfected cells.

15 (I) Comparative costimulatory function of poB7-2 with human B7-1 in the context of MHC restriction.
(II) Flow cytometric analysis of specific anti-poB7-2 antibodies in the sera of immunised mice.
(III) Generation of specific anti-poB7-2 monoclonal antibodies.

20 (I) Comparative *in vitro* analysis can be performed to determine the costimulatory function of poB7-2 or poB7-1 in the context of the human MHC class II molecule HLA-DR1, with that of human B7-1 or B7-2 in the context of DR1, in proliferation assays with human or porcine responders.

25 (II) Transfected P815 cells are crucial reagents for the detection of porcine anti-B7-2 antibody in the sera of immunised mice which have undergone the chimeric peptide immunisation regimen. Flow cytometric analysis with control or poB7-2 -transfected P815 cells, will reflect the specificity of sera for B7-2. Preliminary studies with C57BL-6

mice immunised with a pool of all nine B7-2 peptides have demonstrated the preferential binding of B7-2 peptide sera to porcine B7-2 transfected P815 cells (Figure 11).

(III) Mab with specificity for poB7-2 will be generated by immunisation of Balb/c mice with poB7-2 expressing P815 cells. The spleens from immunised mice will be fused with the NS0 fusion partner and successful fusion's selected by virtue of HAT selection. Flow cytometric staining of poB7-2 P815 transfectants with culture supernatants will enable the identification of MAb secreting cells. Cells will be grown in culture and the medium harvested for antibody purification by passage over Protein G following ammonium

10 sulphate precipitation.

MAb with specificity for B7-1 and CD40 will be generated using the same protocol once the appropriate clones have been obtained. These MAb will provide valuable reagents for further characterising the expression of CS molecules on relevant porcine tissues.

15

5.3 Design and synthesis of poB7-2/OVA chimeric peptide constructs

Nine different peptides derived from the sequence of poB7-2 were initially selected for synthesis. Repeated batches of different peptides will be synthesised until successful molecules are obtained. Porcine B7-2 peptides, 6-22mer in size, were selected as

20 determined by the predicted size of a B cell epitope. Peptides were selected for synthesis in combination with a T cell epitope OVA 323-339. B7-2 peptides were selected on the basis of 3D computer modelling (in collaboration with Paul Travers) and on the basis of predicted antigenicity and hydrophilicity using the SeqAid II computer software package. All of the nine peptides reflect linear epitopes. The positions of the nine peptides in the 25 cloned poB7-2 sequence are indicated (Figure 12). Synthetic peptide sequences are detailed in Table 1

Table 1

Peptide Name	Peptide Sequence	Position
Peptide 1	ISQAVHAAHAEINEAGRSFDQATWTLR	81-90
Peptide 2	ISQAVHAAHAEINEAGRLPCHFTNSQ	32-40
Peptide 3	ISQAVHAAHAEINEAGRKGPHGLVPIHQMS	109-121
Peptide 4	ISQAVHAAHAEINEAGRGLVPIHQMS	113-121
Peptide 5	ISQAVHAAHAEINEAGRQVKDKGSYQC	94-104
Peptide 6	ISQAVHAAHAEINEAGRCSSTQGYPEPQR	151-162
Peptide 8	ISQAVHAAHAEINEAGRKSQAYFNETGEL	21-32
Peptide 9	ISQAVHAAHAEINEAGRASLKSQAYFNET	17-29
Peptide 10	ISQAVHAAHAEINEAGRYMGRTSFDQATWT	76-88
Ova Peptide	ISQAVHAAHAEINEAGR	323-339

5 The peptide sequences and amino acid positions for peptides 1-10 relate to the position of the B7-2 peptide sequence within porcine B7-2. The amino acid position for the ova sequence is only indicated for the Ova peptide. A 17 amino acid peptide from chicken egg albumin (ovalbumin) was selected as the T cell epitope, OVA323-339 (ISQAVHAAHAEINEAGR). This epitope was selected on the basis of published reports
10 for the generation of a H-2^b restricted T cell response (76,77). We have demonstrated the ability of C57BL-6 mice (H-2^b haplotype) to mount a proliferative response to both the native molecule and to the OVA 323-339 peptide following immunisation with whole ovalbumin (Figure 13). Peptides were generated on a peptide synthesiser (Genosys) and crude peptides were purified by HPLC to greater than 70% purity. Sera from OVA
15 control immunised mice should ideally not recognise the 323-339 sequence, indicating that the T cell epitope is devoid of B cell determinants.

5.4 Tolerance induction

5.4.1 *In vivo* tolerance induction strategy

20 C57BL-6 mice will be immunised with whole ovalbumin in CFA, followed by either control peptide (OVA peptide) or test peptides (OVA-B7-2 constructs) for three weekly immunisations. Blood will be collected following sacrifice and sera prepared using a

(iii) Following the successful cloning of all three costimulatory molecules, a combined strategy will be employed to block all three CS molecules by immunisation with appropriate peptides. It is predicted that blocking all three CS molecules will be sufficient to inhibit T cell mediated destruction of the graft by the direct pathway resulting in 5 prolongation of islet graft survival. The tolerance induction strategy detailed in this application is directed against the direct xenorecognition pathway. Thus, if islet survival is to be significantly enhanced above that of controls, other additional strategies may be necessary to target the indirect pathway.

- 10 The results obtained with B7-2 to date, demonstrate the ability of synthetic B7-2 peptides conjugated to a known T cell helper epitope to generate the production of anti-porcine B7-2 antibody *in vivo*. These antibodies if directed towards the binding site between B7 isoforms and CD28, in association with antibodies directed against CD40-CD40L will block the costimulation of human T cells with direct anti-pig xenoreactivity thereby 15 prolonging islet survival in a xenotransplantation context.

Having established the suitability of such an approach in a pig islet to mouse *in vivo* model, studies would progress to pig to primate transplantation systems prior to clinical trials.

- 20
- 5.5 **Adaptations for clinical use of these strategies**
For clinical applicability the following requirements will be necessary:
 - (I) selection of a suitable T cell epitope to replace OVA. One candidate molecule is tetanus toxoid (TT) which is a widely used antigen for use in human immunisation 25 strategies (68,86). The prior immunisations of most adults with TT is an additional benefit to this strategy as memory T cells are already present in the circulation.
 - (ii) An efficient and rapid screening method will be required to detect the presence of anti-donor (pig) B7-2 antibodies in the absence of a specific B7-2 directed T cell response generated by the recipient which would accelerate graft rejection.

6. SUMMARY OF SPECIFIC EMBODIMENTS

The above examples relate to a novel strategy to inhibit costimulation by porcine cells of human T cells with direct anti-pig xenoreactivity. This is of particular importance in the 5 context of xenotransplantation of porcine organs due to the expression of costimulatory molecules on porcine endothelial, as well as bone marrow-derived antigen presenting cells.

Recipients will be immunised with hybrid synthetic peptides comprising a T cell epitope 10 conjugated to sequences of the porcine costimulatory molecules, CD80, CD86 and CD40. Peptides will be selected that induce antibodies specific for regions of the costimulatory molecules involved in binding to their counter-receptors on human cells (CD28 and CD154), and therefore capable of blocking the delivery of costimulation. Once the antibody response has been induced, the transplanted organ will recall this response due 15 to the expression of the costimulatory molecules, thereby sustaining this response, and providing an endogenous mechanism of costimulatory blockade.

standard technique. Presence of specific mouse anti-porcine B7-2 IgG and/or IgM Ab will be detected by one of two strategies.

Peptide ELISAs will be used to screen for the presence of anti-peptide antibody in the sera. Peptides are coated to plates by virtue of aldehyde linkages to allow free access of Ab to the peptide (78). Plates will be coated with individual peptides or the ova control peptide to enable the identification of specific peptides of interest. To detect reactivity of sera with the native B7-2 molecule expressed on the surface of PoB7-2 transfected P815 cells, flow cytometry will be performed following surface staining. Having identified a candidate peptide of interest (peptide ELISA positive and recognising native B7-2) the sera will be used to try to inhibit *in vitro* T cell proliferative responses. This will determine whether the antibody is a blocking antibody. *In vivo* studies will then be performed using the islet transplant system. Antibodies which recognise the native molecule but fail to block a proliferative response will still be useful polyclonal antibody reagents.

To date, initial immunisations involved two groups of mice, one received a pool of all nine B7-2 peptides, and one receiving ova control peptide. The harvested sera were screened by peptide ELISA (Figure 14) which enabled the identification of potential peptides of interest. Peptides 2, 4 and 6 clearly demonstrate preferential binding to B7 peptide than to ova control. The sera has also demonstrated enhanced binding to poB7-2 transfected cells (Figure 11). Peptide 4 was selected as a candidate peptide and used in a subsequent immunisation protocol. Immunisation with peptide 4 alone clearly produced a significant level of IgG with specificity for peptide 4 in the sera of immunised mice (Figure 15). The specificity of the sera for peptide 4 and not to ova control is demonstrated in Figure 16. The ability of sera from peptide 4 immunised mice to specifically recognise the native porcine B7-2 molecule expressed on the surface of porcine B7-2 transfected P815 cells is illustrated in Figure 17. Untransfected control P815 cells do not stain with the Peptide 4 sera, neither do control or transfected cells incubated with ova peptide sera. Similar protocols will be followed with peptides 2 and

6. These data clearly demonstrate the ability of this technique to generate anti-peptide antibody directed against an amino acid sequence, by virtue of a carrier T cell epitope. An identical strategy will be followed with peptides designed on the basis of porcine CD40 and porcine B7-1 once the DNA sequence encoding these molecules has been 5 elucidated.

5.4.2 Functional assessment; prolongation of pancreatic islet xenograft survival

Islet xenografts being non-vascular are rejected solely by T cell mediated mechanisms (79,80), thereby providing an ideal system to study modulation of T cell mediated 10 reactions. A very clear role for cell mediated rejection of islets has been demonstrated and is reported to be greater than the comparable alloresponse (80). Transplantation of porcine pancreatic islets to mice is an established procedure, which is well documented in the literature (80-83). Preliminary studies within this laboratory have demonstrated a decrease in hyperglycaemia (Figure 18) following transplantation of pancreatic islets 15 from large white pigs under the kidney capsule of C57BL-6 mice rendered diabetic by intraperitoneal administration of streptozotocin. Further optimisation of the isolation procedure (84,85) is required to enable purification of fully functional islets. Transplanted islets usually survive between 6-10 days in the absence of any immunosuppression. Successful modulation of direct T cell mediated xenorejection will be monitored by 20 prolongation of islet survival beyond day 10, with comparison to the appropriate controls.

Plans for investigation include:

- (i) Survival and functional assessment of transplanted islets from control and test mice. Survival of islets ie: graft tolerance, will be determined by reversion to and maintenance 25 of normoglycaemia as monitored by using a Reflolux S blood glucose meter. Prior to islet transplantation mice will follow the tolerance induction strategy detailed above.
- (ii) To determine whether our tolerance induction strategy has induced graft-specific T cell tolerance, identical or third party islets can be transplanted under the contra-lateral kidney capsule.

7. Bibliography

1. Dorling, A. *et al.* Clinical Xenotransplantation. *Lancet.* (1997). 349:867-71.
- 5 2. Cooper, D.K.C. Xenografting: how great is the clinical need. *Xeno.* (1995). 1: 25-26
3. Advisory Group on the Ethics of Xenotransplantation. *Animal Tissue into Humans.* London: Stationery Office, 1997.
- 10 4. Nuffield Council on Bioethics. *Animal-to-human transplants.* London: Nuffield Foundation, 1996.
- 15 5. van Denderen, B.J. *et al.* Combination of decay-accelerating factor expression and alpha 1,3-galactosyltransferase knockout affords added protection from human complement-mediated injury. *Transplantation.* (1997). 64. 882-888.
6. Thompson, C. Humanised pigs hearts boost xenotransplantation. *Lancet* (1995): 346: 766.
- 20 7. Dorling, A. *et al.* Detection of primary direct and indirect human anti-porcine T cell responses using a porcine dendritic cell population. *European Journal of Immunology* (1996): 26: 1378.
- 25 8. Dorling, A. *et al.* Cellular xenoresponses: Observation of significant primary indirect human T cell anti-pig xenoresponses using co-stimulator-deficient or SLA class II-negative porcine stimulators. *Xenotransplantation* (1996): 3: 112.
9. Kirk, AD. *et al.* In-vitro analysis of the human anti-porcine T-cell repertoire. *Transplantation Proceedings.* (1992): 24: 602.
- 30 10. Murray, AG. *et al.* Porcine aortic endothelial cells activate human T cells: Direct presentation of MHC antigens and costimulation by ligands for human CD2 and CD28. *Immunity* (1994): 1: 57.

11. Yamada, K. *et al*. Human anti-porcine xenogeneic T cell response. *The Journal of Immunology*. (1995). 155: 5249-5256.

12. Kumagai-Braesch, M. *et al*. Characteristics of direct and indirect activation of human T cells against allogeneic and porcine xenogeneic cells/peptides. *Xenotransplantation*. (1997). 4 : 85-94.

13. Dorling, A. and Lechler, R.I. The passenger leukocyte, dendritic cell and antigen-presenting cells (APC), In *Transplantation Biology; Cellular and Molecular Aspects*. Eds N. L. Tilney, T. B. Strom and L. C. Paul. Philadelphia: Lippincott-Raven, 1996.

14. Lenschow, D.J. *et al*. Inhibition of transplant rejection following treatment with anti-B7.1 antibodies. *Transplantation*. (1995). 60 : 1171-1178.

15. Bretscher, P. and Cohen, M. A theory of self-nonsel discrimination. *Science* (1970): 169: 1042.

16. Bretscher, P. The two signal theory of lymphocyte activation twenty one years later. *Immunology Today*. (1992). 13 : 74-76.

17. Mueller, D.L. *et al*. Clonal expansion versus functional clonal inactivation : A costimulatory pathway determines the outcome of T cell receptor occupancy. *Annual Reviews of Immunology*. (1989). 7 : 445-480.

18. Mueller, D.L. *et al*. An accessory cell-derived costimulatory signal acts independently of protein kinase C activation to allow T cell proliferation and prevent the induction of unresponsiveness. *The Journal of Immunology*. 142: 2617-2628.

19. Baird, M.A. Evidence that heat-treated antigen-presenting cells induce hyporesponsiveness in allogeneic T cells. *Transplantation*. (1994): 57: 763.

20. Jenkins, M.K. *et al*. Molecular Events in the induction of a non-responsive state in interleukin 2 producing helper T- Lymphocyte clones. *Proceedings of the National Academy of Science USA* (1987): 84: 5409.

21. Inaba, K. and Steinman, RM. Resting and sensitized T lymphocytes exhibit distinct stimulatory (antigen-presenting cell) requirements for growth and lymphokine release. *Journal of Experimental Medicine* (1984): 160: 1717.

5

22. Schwartz, R.H. A cell culture model for T lymphocyte clonal anergy. *Science*. (1990). 248: 1349-1355.

10

23. Lafferty, K.J. *et al.* Immunobiology of tissue transplantation: A return to the passenger leukocyte concept. *Annual Reviews of Immunology*. (1983): 1: 143.

24. Jenkins, M.K. and Schwartz, R.H. Antigen presentation by chemically modified splenocytes induces antigen-specific T cell unresponsiveness *in vivo* and *in vitro*. *The Journal of Experimental Medicine*. (1986). 165: 302-319.

15

25. Schultze, J. *et al.* B7-mediated costimulation and the immune response. *Blood Reviews*. (1996). 10 : 111-127.

20

26. June, C.H. *et al.* The B7 and CD28 receptor families. *Immunology Today* (1994): 15: 321.

27. Freeman, G.J. *et al.* B7, A new member of the Ig Superfamily with unique expression on activation and neoplastic B cells. *Journal of Immunology*. (1989): 143: 25 2714.

28. Freeman, G.J. *et al.* Cloning of B7-2: A CTLA-4 counter receptor that co-stimulates human T cell proliferation. *Science* (1993): 262: 909.

30

29. Azuma, M. *et al.* B70 antigen is a second ligand for CTLA-4 and CD28. *Nature* (1993): 366: 76.

35

30. Linsley, P.S. T-cell antigen CD28 mediates adhesion with B cells by interacting with activation antigen B7/BB1. *Proceedings of the National Academy of Science USA* (1990): 87: 5031.

31. Norton, S.D. *et al.* The CD28 Ligand B7, Enhances IL-2 Production by Providing a Costimulatory Signal to T Cells. *Journal of Immunology* (1992): 149: 1556.

5 32. Galvin, F. *et al.* Murine B7 antigen provides a sufficient costimulatory signal for antigen-specific and MHC-restricted T cell activation. *Journal of Immunology* (1992): 149: 3802.

10 33. Boussiotis, VA. *et al.* Activated human B lymphocytes express three CTLA-4 counterreceptors that costimulate T-cell activation. *Proceedings of the National Academy of Science. U S A* (1993): 90: 11059.

34. vanGool, S.W. CD80, CD86 and CD40 provide accessory signals in a multiple step T cell activation model. (1996). 153: 47-83.

15 35. Tang, A. *et al.* Blockade of CD40-CD40 ligand pathway induces tolerance in murine contact hypersensitivity. *European Journal of Immunology*. (1997). 27: 3143-3150.

36. Grewal, I.S. and Flavell, R.A. The role of CD40 ligand in costimulation and T cell activation. *Immunological Reviews*. (1996). 153: 86-106.

20 37. Sun, H. *et al.* Prevention of chronic rejection in mouse aortic allografts by combined treatment with CTLA4Ig and anti-CD40 ligand monoclonal antibody. *Transplantation*. (1997). 64: 1838-1856.

25 38. Larsen, C.P. *et al.* Longterm acceptance of skin and cardiac allografts after blocking CD40 and CD28 pathways. *Nature*. (1996). 381: 434-441.

39. Wecker, H. and Auchincloss, H. Cellular mechanisms of rejection. *Current Opinion in Immunology*. (1992). 4: 561-566.

30 40. Satake ,M. *et al.* Direct activation of human responder T cells by porcine stimulator cells

leads to T cell proliferation and cytotoxic T cell development. *Xenotransplantation*. (1996). 3: 198-206.

41. Kirk, A.D. *et al.* The human anti-porcine T cell repertoire. In vitro studies of acquired 5 and innate cellular responsiveness. *Transplantation*. (1993). 55 : 924-931.

42. Alter, B. and Bach, F.H. Cellular basis of the proliferative response of human T cells to mouse xenoantigens. *Journal of Experimental Medicine*. (1990). 171: 333-338.

10 43. Baliga, P. *et al.* CTLA4Ig prolongs allograft survival while suppressing cell mediated immunity. *Transplantation* (1994): 58: 1082.

15 44. Turka, LA. T cell activation by the CD28 ligand B7 is required for cardiac allograft rejection *in vivo*. *Proceedings of the National Academy of Science, USA* (1992): 89: 11102.

20 45. Lin, H. *et al.* Long term acceptance of major histocompatibility complex mismatched cardiac allograft induced by CTLA4-Ig plus donor specific transfusion. *Journal of Experimental Medicine* (1993). 178: 1801.

46. Lenschow, DJ. *et al.*. Long term survival of xenogeneic pancreatic islet grafts induced by CTLA4-Ig. *Science*. (1992): 257: 789.

25 47. Lu,L. *et al.* Blockade of the CD40-CD40 ligand pathway potentiates the capacity of donor derived dendritic cell progenitors to induce long-term cardiac allograft survival. *Transplantation*. (1997). 64: 1808-1815

48. Fallarino, F. *et al.* B7-1 engagement of cytotoxic T lymphocyte antigen 4 inhibits T cell activation in the absence of CD28. *Journal of Experimental Medicine*. (1988). 188 : 30 205-210.

49. Freeman, G.J. *et al.* Murine B7-2, an alternative CTLA4 counter-receptor that costimulates T cell proliferation and IL-2 production. *Journal of Experimental Medicine.* (1993). 178: 2185-2192.

5 50. Jenkins, K.M. and Johnson, J.G. Molecules involved in T-cell costimulation. *Current Opinion in Immunology.* (1993) 5 : 361-367.

51. Brunet, J.F. *et al.* A new member of the immunoglobulin superfamily--CTLA-4. *Nature* (1987). 328: 267.

10 52. Lenschow, D.J. *et al.* B7 system of T cell costimulation. *Annual Reviews of Immunology.* (1996). 14 : 233-258.

15 53. Norton, S.D. The CD28 ligand, B7, enhances IL-2 production by providing a costimulatory signal to T cells. *The Journal of Immunology.* (1992). 149 : 1556-1561.

54. Linsley, P.S. *et al.* T cell antigen CD28 mediates adhesion with B cells by interacting with activation antigen B7/BB1. *Proceedings of the National Academy of Science.* (1990). 87 : 5031-5035.

20 55. Krummel, M.F. *et al.* CD28 and CTLA-4 have opposing effects on the response of T cells to stimulation. *Journal of Experimental Medicine* (1995): 182: 459.

25 56. Krummel, M.F. and Allison, J.P. CTLA-4 engagement inhibits IL-2 accumulation and cell cycle progression upon activation of resting T cells. *Journal of Experimental Medicine* (1996): 183: 2533.

57. Walunas, T.L. *et al.* CTLA-4 ligation blocks CD28-dependent T cell activation. *Journal of Experimental Medicine* (1996). 183: 2541.

30 58. Gimmi, C.D. *et al.* Human T-cell clonal anergy is induced by antigen presentation in the absence of B7 costimulation. *Proceedings of the National Academy Science. U S A* (1993): 90: 6586.

59. Waterhouse, P. *et al.* Lymphoproliferative disorders with early lethality in mice deficient in CTLA4. *Science* (1995); 270: 985.

5

60. Maher, S.E. *et al.* Porcine endothelial CD86 is a major costimulator of xenogeneic human T cells. *The Journal of Immunology*. (1996). 157: 3838-3844.

10 61. vanEessen, D. *et al.* CD40 ligand-transduced co-stimulation of T cells in the development of helper function. *Nature*. (1995) 378. 620-623.

62. Larsen, C.P. *et al.* CD40-gp39 interactions play a critical role during allograft rejection. *Transplantation*. (1996). 61: 4-9.

15

63. Larsen, C.P. and Pearson, T.C. The CD40 pathway in allograft rejection, acceptance and tolerance. *Current Opinion in Immunology*. (1997). 9: 641-647.

20 64. Bennet, S.R.M. *et al.* Help for cytotoxic T-cell responses is mediated by CD40 signalling. *Nature*. (1998). 393: 478-480.

65. Schoenberger, S.P. *et al.* T cell help for cytotoxic T lymphocytes is mediated by CD40-CD40L interactions. *Nature*. (1998) 393: 480-483.

25 66. Ridge, J. P. *et al.* A conditioned dendritic cell can be a temporal bridge between a CD4 T helper and a T-killer cell. *Nature*. (1998) 393: 474-478.

67. Tran, H.M. *et al.* Short-term xeno-suppression of the xeno-immune response with mCTLA4-Fc treatment. *Transplantation*. (1997). 4: 222-227

30

68. Lise, L.D. *et al.* Enhanced epitopic responses to a synthetic human malarial peptide by preimmunisation with tetanus toxoid carrier. *Infection and Immunity*. (1987). 55: 2658-2661.

5 69. Dalum, I. *et al.* Breaking of B cell tolerance toward a highly conserved self protein. *The Journal of Immunology*. (1996). 157: 4796-4804.

70. Dalum, I. *et al.* Induction of cross-reactive antibodies against a self-protein by immunisation with a modified self protein containing a foreign T helper epitope.

10 *Molecular Immunology*. (1997). 34: 1113-1120.

71. Sad, S. *et al.* Bypass of carrier induced epitope-specific suppression using a T helper epitope. *Immunology*. (1992). 76: 599-603.

15 72. Sad, S. *et al.* Carrier induced suppression of the antibody response to a "self"-hapten. *Immunology*. (1991). 74: 223-227.

73. Grimaldi, J.C. *et al.* Genomic structure and chromosomal mapping of the murine CD40 gene. *The Journal of Immunology*. (1992). 149: 3921-3926.

20 74. Stamenkovic, I. *et al.* A B lymphocyte activation molecule related to the nerve growth receptor and induced by cytokines in carcinomas. *The EMBO Journal*. (1989). 8: 1403-1410.

25 75. Ramesh, N. *et al.* Chromosomal localisation of the gene for human B-cell antigen CD40. *Somatic Cell and Molecular Genetics*. (1993). 19: 295-298.

76. Shimonkevitz, R. *et al.* Antigen recognition by H-2-restricted T cells. *The Journal of Immunology*. (1984). 133: 2067-2074.

77. Robinson, J. H. *et al*. Palmitic acid conjugation of a protein antigen enhances major histocompatibility complex class II restricted presentation to T cells. *Immunology*. (1992) 76 : 593-598.

5 78. Elma, E.M.G. *et al*. Direct coating of poly(lys) or acetyl-thio-acetyl peptides to polystyrene: The effects in an enzyme-linked immunosorbent assay. *Analytical Biochemistry*. (1997). 248: 117-129.

● 10 79. Wennberg, L. *et al*. Allogeneic and xenogeneic islets are rejected by different and specific mechanisms: A study in rodents using a mixed allogeneic-xenogeneic islet transplantation model. *Xenotransplantation*. (1997). 4 : 228-234.

● 15 80. Mandel, T.E. *et al*. Cellular rejection of fetal pancreas grafts: differences between allo and xenograft rejection. *Xenotransplantation*. (1997) 4: 2-10.

● 20 81. Mandel, T. E. *et al*. Transplantation of organ cultured fetal pig pancreas in non-obese diabetic (NOD) mice and primates (*Macaca fascicularis*). *Xenotransplantation*. (1995) 2: 128-132.

● 25 82. Lu, X. *et al*. Long-term survival of hamster islet xenografts in mice under short course treatment with non depleting versus depleting anti-CD4 monoclonal antibodies. *Xenotransplantation*. (1998). 5 : 154-163.

83. Marchetti, P. *et al*. Automated large-scale isolation, *in vitro* function and 25 xenotransplantation of porcine islets of langerhans. *Transplantation*. (1991). 52: 209-213.

● 20 84. Ricordi, C. *et al*. A method for the mass isolation of islets from the adult pig pancreas. *Diabetes*. (1986) 35: 649-653.

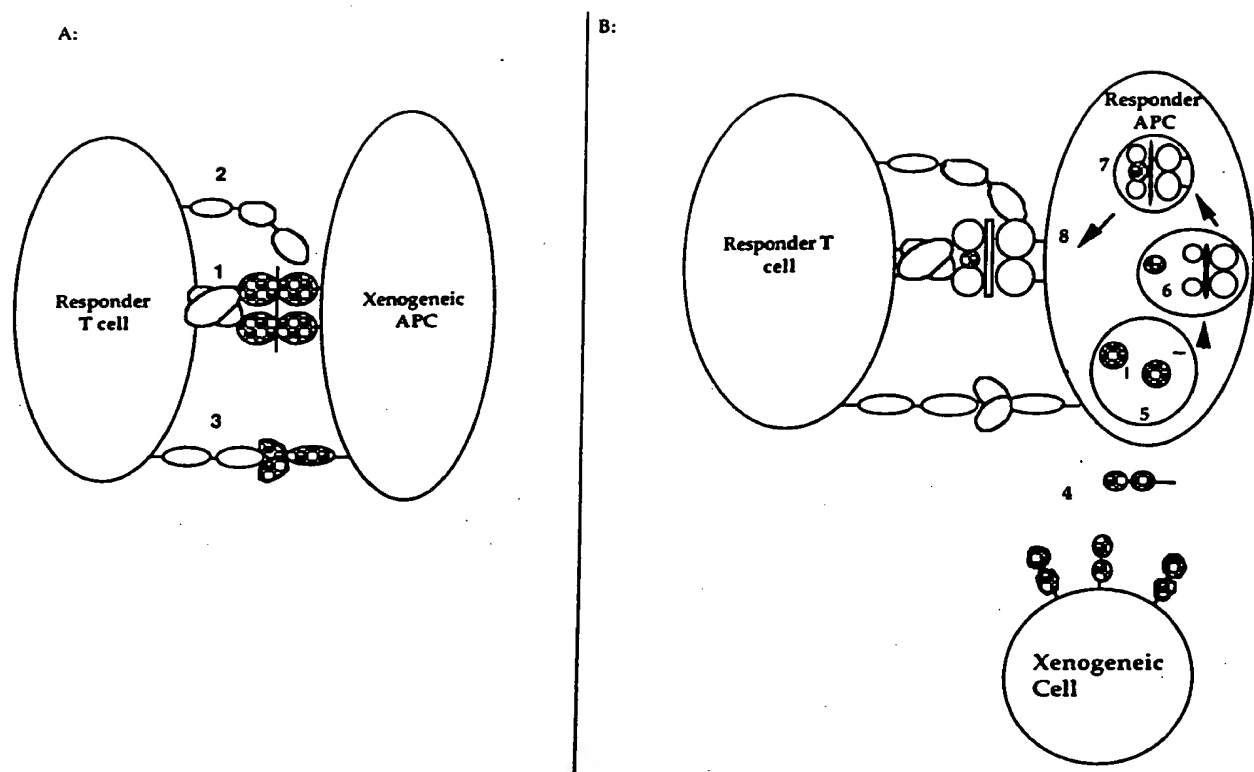
30 85. Ricordi, C. *et al*. Isolation of the elusive pig islet. *Surgery*. (1990). 107: 688-694.

86. Etlinger, H.M. *et al.* Use of prior vaccinations for the development of new vaccines. Science. (1990). 249: 423-425.

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p15700

Figure 1



A: Diagrammatic representation of direct xenorecognition.

The types of molecular interactions necessary for efficient direct xenorecognition are numbered 1 - 3.

1 Cognate interaction between TCR on responder T cell and MHC molecules on xenogeneic antigen presenting cells.

2 Non cognate interaction between co-receptors CD4 and membrane proximal domains of MHC class II, and CD8 and $\alpha 3$ domains of MHC class I.

3 Non cognate interactions between accessory and costimulatory molecules. Important interactions are between B7 family (APC) and CD28 (T), LFA-3 (APC) and CD2 (T), and ICAM-1 (APC) and LFA-1 (T)

B: Diagrammatic representation of indirect xenorecognition

Xenoantigens (4), released by xenogeneic cells, are taken up and processed (5) into peptide fragments by specialised antigen presenting cells (6) before binding to MHC class II molecules (7) and display on the cell surface (8) for presentation to xenospecific self-class II MHC-restricted T cells.

Figure 1: Diagrammatic comparison of direct and indirect xenorecognition pathways.

GCATGGATCCATGGGACTGAGTAACATTCTTTG
 1 **ATGGGACTGAGTAACATTCTTTG**ATGGTCCTCCT
 39 GCTCTCTGGTGCCTCCTTGAAAAAGTCAGGCATATTCAATGAGA
 86 CTGGAGAACTGCCGTGCCATTACAAACTCGCAGAACCTAACGCTG
 133 GATGA**G**CTGGTCATATTGGCAGGACCAGGATAACCTGGTTCTCTA
 181 CGAGCTATAACGAGGCCAAGAGAACGCCTCATAATGTTAATTCCAAG
 227 TATATGGGTCGCACAAGCTTGACCAAGGCCACCTGGACCCCTGAGACT
 274 CCACAACTGTTCAAATCAAGGACAAGGGCTCATATCAATGTT**C**ATC
 321 CATCATAAAGGGCCGCATGGACTTGTCTATCCACCAAGATGAGTT
 368 TGACCTATCATTGCTGCTAACCTCAGTCACCTGAAATAAACCTAC
 415 TTACTAATCACACAGAAAATTCTGTCTAAATTGACCTGCTCATCT
 462 ACACAAAGGCTACCCAGAACCCAGAGGATGTATATGTTGCTAAATA
 509 CGAAGAA**F**TCACCAACTGAGCATGATGCTGACATGAAGAAATCTCA
 556 AAATAACATCACGGAACTCTACAATGTATCAATCAGGGTGTCTCTT
 602 CCCATCCCTCCCGAGACAAATGTGAGCATCGTCTGTGCTGCAACTT
 649 GAGCCAAGCAAGACACTGCTTTCTCCCTACCTGTAATATAGATGC
 696 AAAGCCACCTGTGCAACCCCTGTCCCAGACCACATCCTCTGGATTGC
 743 AGCTCTACTTGTAAACAGTGGTCGTTGTGTGGATGGTGTCTTTGT
 790 AACACTAAGGAAAAGGAAGAAGAAGAACGAGCCTGGCCCTCTAATGA
 837 ATGTGGTAAACCCATCAAAATGAACAGGAAGGGCGAGTGAACAAAC
 884 TAAGAACAGAGCAGAACGATCTGATGATGCCAGTGT
 931 GATGTTAATATTTAAAGACAGCCTCAGATGACAACAGTACTACAG
 GACAACAGTACTACAG
 978 **ATTTTAAATTAAAGAGTAAACTCC**
 - **ATTTTAAAGTCGACATGC**

Figure 2: Position of 5' and 3' primers (highlighted in bold type) with respect to the published coding sequence of porcine CD86. The underlined sequences ATG and TAA represent the start and stop codons respectively.

1 CACCGCGGTG CGGCCGCTCT AGAACTAGTG GATCCATGGG ACTGAGTAAC
51 ATTCTCTTG GGATGGTCCT CCTGCTCTCT GGTGCTGCCCT CCTTGAAAAG
101 TCAGGCATAT TTCAATGAGA CTGGAGAACT GCCGTGCCAT TTACAAACT
151 CGCAGAACCT AAGCCTGGAT GAGCTGGTCA TATTTGGCA GGACCAGGAT
201 AACCTGGTTC TCTACGAGCT ATACCGAGGC CAAGAGAAC CTCATAATGT
251 TAATTCCAAG TATATGGGTC GCACAAAGCTT TGACCAGGCC ACCTGGACCC
301 TGAGACTCCA CAACGTTCAA ATCAAGGACA AGGGCTCATA TCAATGTTTC
351 ATCCATCATA AAGGGCCGCA TGGACTTGTT CCTATCCACC AGATGAGTTC
401 TGACCTATCA GTGCTTGCTA ACTTCAGTCA ACCTGAAATA AACCTACTTA
451 CTAATCACAC AGAAAATTCT GTCATAAATT TGACCTGCTC ATCTACACAA
501 GGCTACCCAG AACCCCAGAG GATGTATATG TTGCTAAATA CGAAGAATTC
551 AACCACTGAG CATGATGCTG ACATGAAGAA ATCTAAAAT AACATCACGG
601 AACTCTACAA TGTATCAATC AGGGTGTCTC TTCCCATCCC TCCCGAGACA
651 AATGTGAGCA TCGTCTGTGT CCTGCAACTT GAGCCAAGCA AGACACTGCT
701 TTTCTCCCTA CCTTGTAATA TAGATGAAA GCCACCTGTG CAACCCCCTG
751 TCCCAGACCA CATCCTCTGG ATTGCAGCTC TACTTGTAAAC AGTGGTCGTT
801 GTGTGTGGGA TGGTGTCCCT TGTAAACACTA AGAAAAAGGA AGAAGAAGCA
851 GCCTGGCCCC TCTAATGAAT GTGGTGAAAC CATAAAATG AACAGGAAGG
901 CGAGTGAACA AACTAAGAAC AGAGCAGAAC TCCATGAACG ATCTGATGAT
951 GCCCAGTGTG ATGTTAATAT TTTAAAGACA GCCTCAGATG ACAACAGTAC
1001 TACAGATTT TAAGTCGACC TCGAGGGGGG GCCCGGTACC AGCTTTGTT

Figure 3: Nucleotide sequence of CD86(i) obtained by RT-PCR amplification of cDNA extracted from a transformed porcine endothelial cell line A8.

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Figure 4: Comparison of the nucleotide sequence of CD86(i) with the published sequence for porcine CD86.

Figure 4:

ATGGGACTGAGTAACATTCTCTTGATGGTCCTCTGCTCTCTGG
 CACCGCGGTGCGGCCGCTCTAGAACTAGTGGATCCATGGGACTGAGTAACATTCTCTTGATGGTCCTCTGCTCTCTGG
 | 10 | 20 | 30 | 40 | 50 | 60 | 70 | 80 |
 TGCTGCCTCTTGAAAAGTCAGGCATATTCATGAGACTGGAGAACTGCCGTGCCATTAACTCGCAGAACCTAACGC
 TGCTGCCTCTTGAAAAGTCAGGCATATTCATGAGACTGGAGAACTGCCGTGCCATTAACTCGCAGAACCTAACGC
 | 50 | 60 | 70 | 80 | 90 | 100 | 110 | 120 | 130 | 140 | 150 | 160 |
 CTGGATGAGCTGGTCATATTTGGCAGGACCAGGATAACCTGGTTCTCTACGAGCTATACCGAGGCCAAGAGAACCTCATA
 CTGGATGAGCTGGTCATATTTGGCAGGACCAGGATAACCTGGTTCTCTACGAGCTATACCGAGGCCAAGAGAACCTCATA
 | 130 | 140 | 150 | 160 | 170 | 180 | 190 | 200 | 210 |
 ATGTTAATTCCAAGTATATGGGTCGCACAAGCTTGGACCAAGGCCACCTGGACCCCTGAGACTCCACAACGTTCAAATCAAGGA
 ATGTTAATTCCAAGTATATGGGTCGCACAAGCTTGGACCAAGGCCACCTGGACCCCTGAGACTCCACAACGTTCAAATCAAGGA
 | 220 | 230 | 240 | 250 | 260 | 270 | 280 | 290 | 300 | 310 | 320 |
 CAAGGGCTCATATCAATGTTTCCATCATAAAGGGCCGATGGACTTGGCTTCTATCCACCAAGATGAGTTCTGACCTATCA
 CAAGGGCTCATATCAATGTTTCCATCATAAAGGGCCGATGGACTTGGCTTCTATCCACCAAGATGAGTTCTGACCTATCA
 | 300 | 310 | 320 | 330 | 340 | 350 | 360 | 370 | 380 | 390 | 400 | 410 |
 TTGCTTGCTAACCTCAGTCAACCTGAAATAAACCTACTTACTAACACACAGAAAATTCTGTCATAAATTGACCTGCTCAT
 GTGCTTGCTAACCTCAGTCAACCTGAAATAAACCTACTTACTAACACACAGAAAATTCTGTCATAAATTGACCTGCTCAT
 | 380 | 390 | 400 | 410 | 420 | 430 | 440 | 450 | 460 | 470 | 480 | 490 |
 CTACACAGGCTACCCAGAACCCAGAGGATGTATATGTTGCTAAATACGAAGAACCAACTGAGCATGATGCTGACAT
 CTACACAGGCTACCCAGAACCCAGAGGATGTATATGTTGCTAAATACGAAGAACCAACTGAGCATGATGCTGACAT
 | 460 | 470 | 480 | 490 | 500 | 510 | 520 | 530 | 540 | 550 | 560 | 570 |

Contig	ACCATGGGACTGAGTAACATTCTCTTGTGATGGTCTCCTGCTCTCT
Murine B7-2	- <u>CCAT</u> GGGACTGAGTAACATTCTCTTGGGATGGCCTCCTGCTCTCT
Porcine CD68(i)	ACCAT <u>GGG</u> CTTGGCAATCCTATCTTGTGACAGTCTGCTGATCTCA
Human B7.2	ACT <u>AT</u> GGGACTGAGTAACATTCTCTTGTGATGGCCTCCTGCTCTCT


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GGTCTGCTTCBTGAAGABTCAGCTTATTCAATGAGACTGCAGAHCTGCCGTGCCAATTAA
GGTCTGCTCCTCTGAAAGTCAGGCATATTCAATGAGACTGGAGAACTGCCGTGCCAATTAA
GATGCTTTCCTGGAGACGCAAGCTTATTCAATGGGACTGCATATCTGCCGTGCCAATTAA
GGTCTGCTCCTCTGAAGATTCAAGCTTATTCAATGAGACTGCAGACCTGCCATGCCAATTAA

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CAAACCTCAAAACCTAACGCTGAGTGAGCTGGTAGTATTGGCAGGACCAGGAAAACCTGGT
CAAACCTCGAGAACCTAACGCTGGATGAGCTGGTCATATTGGCAGGACCAGGATAACCTGGT
CAAAGGCTCAAAACATAAACGCTGAGTGAGCTGGTAGTATTGGCAGGACCAGCAGGAAAACCTGGT
CAAACCTCAAAACCAAAACGCTGAGTGAGCTAGTAGTATTGGCAGGACCAGGAAAACCTGGT

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TCTGTACGAGCTATACTTACCGAAAGAGAAACTTGATAGTTAATTCAAGTATATGGGCCGC
TCTCTACGAGCTATAACCGAGGCCAACGCTCATAATGTTAATTCAAGTATATGGGTGCG
TCTGTACGAGCACTATTGGGCACAGAGAAACTTGATAGTGTGAATGCCAGTACCTGGGCCGC
TCTGAATGAGGTATACTTACCGAAAGAGAAATTGACAGTGTCAATTCAAGTATATGGGCCGC

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ACAAGCTTGTGACHVGGACAVCTGGACCCCTGAGACTTCACAATGTTAGATCAAGGACAAGGGCT
ACAAGCTTGTGACCAAGGCCACCTGGACCCCTGAGACTCCACAACGTTAACTCAAGGACAAGGGCT
ACGAGCTTGTGACAGGAACAACGGACTCTACGACTTCACAATGTTAGATCAAGGACATGGGCT
ACAAGCTTGTGACAGTGGACCCCTGAGACTTCACAATCTCAGATCAAGGACAAGGGCT

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CGTATCAATGTTCATCCATCAHAAAUVGCCACAGGAHTDATTCCBCATCCACCAAGATGADTT
CATATCAATGTTCATCCATCAAAAGGGCCGATGGACTTGTCTATCCACCAAGATGAGTT
CGTATGATTGTTTATACAAAAAAAGCCACCCACAGGATCAATTATCCTCCAACAGACATTAAC
TGTATCAATGTATCATCCATCACAAAAAGGCCACAGGAATGATTGCGATCCACCAAGATGAATT

```



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TGAACGTGTCAGTGCTTGCTAACCTCAGTCACCTGAAATAAAACTAVTHCTAATVTAACAGAA
TGACCTATCAGTGCTTGCTAACCTCAGTCACCTGAAATAAAACCTACTTAATCACACAGAA
AGAACTGTGTCAGTGATGCCAACCTCAGTGAAACCTGAAATAAAACTGGCTCAGAATGTAACAGGA
TGAACGTGTCAGTGCTTGCTAACCTCAGTCACCTGAAATAAGTACCAATTCTAATATAACAGAA

```

Figure 5: Comparison of CD86(i) with published sequences for murine and human CD86. Sequence continues overleaf.

Con | AATTCTGDCATAAATTGACCTGCTCATCTAACAAGGTTACCCAGAACCTAAGAAGATGTATD
 Murine B7.2 | AATTCTGTCATAAATTGACCTGCTCATCTACACAAGGCTACCCAGAACCCAGAGGATGTATA
 Porcine CD68(i) | AATTCTGGCATAAATTGACCTGCACTAAGCAAGGTACCCGAAACCTAAGAAGATGTATT
 Human B7.2 | AATGTGTACATAAATTGACCTGCTCATCTACACGGTTACCCAGAACCTAAGAAGATGAGTG

|||||
 TTTTGCTAAVTACNAAGAATTCAACTAHTGAGTATGATGVTAACTGCAGAAATCTCAAGATAA
 TGTGCTAAATACGAAGAATTCAACCACTGAGCATGATGCTGACATGAAGAAATCTCAAATAA
 TTCTGATAACT-----AATTCAACTAATGAGTATGGTATAACATGCAGATATCACAAGATAA
 TTTTGCTAAGAACCAAGAATTCAACTATCGAGTATGATGGTATTATGCAGAAATCTCAAGATAA

|||||
 TGTCACAGAACTGTACAATGTHTCATCAGCBTGTCTTTCATTCCCTGATGDTACGAGNNAT
 CATCACGGAACCTACAATGTATCAATCAGGGTGTCTTCCCATCCCTCCGAGACAA---AT
 TGTCACAGAACTGTTCACTGAGTATCTCAAACAGCCTCTCTTTCATTCCCGATGGTGTGGCAT
 TGTCACAGAACTGTACGACGTTCCATCAGCTGTCTGTTCATTCCTGATGTTACGAGCAAT

|||||
 ATGACCACATCGTCTGTGTTCTGGAAACTGAGNCANAAGACNCNGCTTTTCTCHHACCTTCA
 GTGAGCATCGTCTGTGTCCTGCAACTTGAGCCAAGCAAGACACTGCTTTTCTCCCTACCTGTA
 ATGACCGTTGTGTGTTCTGGAAACGGAGTCATGAAGA-----TTTCCCTCAAACCTCTCA
 ATGACCACATCTCTGTATTCTGGAAACTGA-----CAAGACGCGGCTTTATCTCACCTTCT

|||||
 ATATAGATCHAGAGBHHCTNNCAACCTCCTNNCCAGACCACATBCNNTGGATTACAGCTBT
 ATATAGATGCAAAGCCACCTGTGCAACCCCTGTCCTCAGACCACATCCTCTGGATTGAGCTCT
 ATTTCACTCAAGAGTTTC-----ATCTCCTCAAACGTATTGGAAG---GAGATTACAGCTTC
 CTATAGAGCTTGAGGACCCT---CAGCCTCC---CCCAGACCACATTCTGGATTACAGCTGT

|||||
 ACTTNNAAACAGTGGTCVTTVTVTGTGTGATGGTGTCTNTVTAATTCTATGGAAANNAAGAAG
 ACTTGTAAACAGTGGTCGTTGTGTGGATGGTGTCTTGTAAACACTAAGGAAA---AGGAAG
 AGTT---ACTGTGGCCCTCCTCTTGTGATGCTGCTC---ATCATIGTATG---TCACAAGAAG
 ACTTCCAACAG---TTATTATATGTGTGATGGTTCTGCTAATTCTATGGAAATGGAAGAAG

|||||
 AAGAACGCAGCTVCAVCTTAATAATGTGGNNNAACCAHAAAATGGAGAGGGANGNGAGTG
 AAGAACGCAGCTGGCCCTCTAATGAATGTGGTGAACCATCAAATGAACAGGAAGGCAGTG
 CCGAACAGCCTAGCAGGCCAGCAA-----CACAGCCTCTAAGTTAGAGCGGGA---TAGT
 AAGAACGCGCTCGCAACTCTTATAATGTGG---AACCAACACAATGGAGAGGGAAAGAGAGTG

|||||
 AACANACTAAGAACAGAGAAAAANTCCATNNACCTGAAVGATCTGATGAAGCCCAGNGTNTNT
 AACAAACTAAGAACAGAGCAGAACAGTCCAT-----GAACGATCTGATGATGCCAGTGTGATGT
 AACG---CTG---ACAGAGAGA---CTATCAACCTGAAGGAACCT---TGAACCCCA-----
 AACAGACCAAGAAAAGAGAAAAATCCATATACTGAAAGATCTGATGAAGCCCAGCGTGTGTTT

|||||
 TAANADTTNNAAAGACAGCTTCANNNGACAAAAGTNNTACANNTTTAADTNAGAGTNAAGNN
 TAATATTTAAAGACAGCCTCAGATGACAACAGTACTACAGATTTTAAGT-----
 ---AATT-----GCTCA---GCAAAA-----CCAAATGCAGAGTGAAG---
 TAAAAGTTGAGACATCTCATGCGACAAAAGTGTACATGTTTAATTAAAGAGTAAAGCC

9/26

Contig	10	20	30	40	50	
Murine CD86	
Porcine CD86(i)	MDPRC-----	TMGLAILIFVTVLLISDAVS	ETQAYFNGTAYLPCPFTKAQNI			
Human CD86	--PRCGRSRTSGSMGLS	NILFGM	VLLLSGAASLKSQAYFNETGELPCHFTNSQL			
Porcine CD86	-----	MGLSNILFVMA	LLSGAAPLKIQAYFNETADLPCQFANSQ	NQ		
	-----	MGLSNILFV	MVLLLSGAASLKSQAYFNETGELPCHFTNSQL			
	60	70	80	90	100	110

	SLSELVVFWQDQQK	LVLYEHYLGTEKLD	SVNAKYLGR	TSFDRNNW	TLLRLHN	VQIK
	DELVIFWQDQDN	LVLYEHYLGKE	TSFDRNNW	TLLRLHN	VQIK	
	DELVIFWQDQEN	LVLYEHYLGKEKF	TSFDRNNW	TLLRLHN	VQIK	
	DELVIFWQDQDN	LVLYEHYLGKEKF	TSFDRNNW	TLLRLHN	VQIK	
	120	130	140	150	160	
	
	DMGSYDCF	IQKKPPTGSI	ILOQTLTE	LSVIANFSE	PEI	KLAQNV
	DKGSYQCF	IHHKGPHGLVPI	HQMSSDL	SVLANFSQ	PEI	NLTCT
	DKGLYQCII	IHHKGPHGLVPI	HQMSSDL	SVLANFSQ	PEI	LNHTEN
	DKGSYQCF	IHHKGPHGLVPI	HQMSSDL	SVLANFSQ	PEI	LNHTEN
	170	180	190	200	210	220
	
	SKQGHPKPK	MYFLIT--NSTNEY	GDNMQ	ISQDNV	TELF	FSISNSL
	SKQGHPKPK	MYFLIT--NSTNEY	GDNMQ	ISQDNV	TELF	FSISNSL
	SKQGHPKPK	MYFLIT--NSTNEY	GDNMQ	ISQDNV	TELF	FSISNSL
	230	240	250	260	270	
	
	MTIVCV	LETESMKISSKPLNFT	QEFPSP-----	QTYW-KEITAS	VTALLVM	
	MTIVCV	LETESMKISSKPLNFT	QEFPSP-----	QTYW-KEITAS	VTALLVM	
	MTIVCV	LETESMKISSKPLNFT	QEFPSP-----	QTYW-KEITAS	VTALLVM	
	280	290	300	310	320	330
	
	LLIIVCHKKP	NQPSRPSN--TAS	KLERDSNAD--RETINL	--KELEPQ	IASA	
	LLIIVCHKKP	NQPSRPSN--TAS	KLERDSNAD--RETINL	--KELEPQ	IASA	
	LLIIVCHKKP	NQPSRPSN--TAS	KLERDSNAD--RETINL	--KELEPQ	IASA	
	340	350				
				
	KPNAE	KTASDDN	STTDFXVDLEGPGTSFC			
	KTSSCD	KSDTCF				
	KTASDDN	STT--DFXLKSKL				

Figure 6: Predicted amino acid sequence for CD86(i) compared with those for pig, human and mice.

Figure 7: Position of 5' and 3' internal and external porcine B7-1 primers with respect to human and murine B7-1 nucleotide sequences. Primer sequences are underlined and labelled as follows. Internal primers (A) and external primers (B).

10 20 30 40 50 60 70 80
 CCAAAGAAAAAGTGATTGTCATTGCTTATAGACTGTAAGAAGAGAACATCTCAGAAGTGGAGTCTTACCCCTGAAATCAAA
 GAGTTTTATAACCTCAATAGACT
 10 20

90 100 110 120 130 140 150 160
 GGATTTAAAGAAAAAGTGGAAATTTCAGCAAGCTGTGAAACTAAATCCACAAACCTTGGAGACCCAGGAACACCCCTCC
 CTTACTAGTTCTCTTTTCAGGTTGTGAAACTCAACCTTCAAAGACACTCTGTTCCATTCTGTGGACTAATAGGATCATC
 30 40 50 60 70 80 90 100

170 180 190 200 210 220 230 240
 AATCTCTGTGTGTTGTAAACATCACTGGAGGGTCTCTACGTGAGCAATTGGATTGTCATCAGCCCTGCCTGTTGCAC
 TTTAGCATCTGCCGGGTGGATGCCATCCAGGTTCTTTCTACATCTCTGTGATTTGTGAGCCTAGGAGGTGCC
 110 120 130 140 150 160 170 180

250 260 270 280 290 300 310 320
 CTGGGAAGTGCCTGGTCTTACTTGGGTCCAATTGGTGGCTTCACCTTGACCCCTAACGATCTGAAGCCATGGGCCACAC
 TAAGCTCCATTGGCTCTAGATTCTGGCTTCCCCATCATGTTCTCCAAAGCATTGAAGCTATGGCTTGAATTGTCAGTT
 190 200 210 220 230 240 250 260

330 340 350 360 370 380 390 400 410
 ACGGAGGCAGGGAACATCACCATCCAAGTGTCCATACCTCAATTCTTCAAGCTCTGGTGTGGCTGGCTTTCTCACCTC
 GATGCAGGATAACCACTCCTCAAGTTCCATGTCCAAGGCTCATTCTCTTTGTGCTGCTGATTGCTTCAACAGTG
 270 280 290 300 310 320 330 340 350

420 430 440 450 460 470 480 490
 TGTTCAAGGTATCCACGTGACCAAGGAAGTGAAGAAGTGGCAACGCTGTGTCCTGTGGTCACAATGTTCTGTTGAAGAGC
 TCTTCAGATGTGATGAACAACTGTCCAAGTCAGTGAAAGATAAGGTATTGCTGCCCTGCCGTTACAACCTCCTCATGAAG
 360 370 380 390 400 410 420 430

500 510 520 530 540 550 560 570
 TGGCACAAACTCGCATCTACTGGCAAAAGGAGAAGAAAATGGTGTGACTATGATGTCCTGGGACATGAATATATGGCCCGA
 ATGAGTCTGAAGAGCCGAATCTACTGGCAAAACATGACAAAGTGGTGTGTCATTGCTGGAAACTAAAAGTGTGGCC
 440 450 460 470 480 490 500 510

580 590 600 610 620 630 640 650
 GTACAAGAACCGGACCATTTGATATCACTAATAACCTCTCATTGTGATCCTGGCTCTGGCCCATCTGACGAGGGCACA
 CGAGTATAAGAACCGGACTTTATATGACAACACTACCTACTCTTATCATCCTGGCCTGGTCTTCAGACCGGGCACA
 520 530 540 550 560 570 580 590

660 670 680 690 700 710 720 730
 TACGAGTGTGTTCTGAAGTATGAAAAAGACGCTTCAAGCGGGAACACCTGGCTGAAGTGACGTTATCAGTCAGCTG
 TACAGCTGTGTCGTTCAAAAGAAGGAAAGAGGAACGTATGAAGTTAACACTTGGCTTAGTAAAGTTGTCCATCAAAGCTG
 600 610 620 630 640 650 660 670

740 750 760 770 780 790 800 810 820
 ACTTCCCTACACCTAGTATATCTGACTTGAATTCACCTCTAAATTAGAAGGATAATTGCTCAACCTCTGGAGGTTT
 ACTTCTCTACCCCCAACATAACTGAGTCTGGAAACCCATCTGCAGACACTAAAGGATTACCTGCTTGTCTCCGGGGTTT
 680 690 700 710 720 730 740 750 760

830 840 850 860 870 880 890 900
 TCCAGAGCCTCACCTCCTGGTGGAAAATGGAGAAGAATTAAATGCCATCAACACAAAGTTCCCAAGATCCTGAAACT
 CCCAAAGCCTCGCTCTTGGTGGAAAATGGAGAAGAATTACCTGGCATCAATACGACAATTCCCAGGATCCTGAATCT
 770 780 790 800 810 820 830 840

910 920 930 940 950 960 970 980
 AGCTCTATGCTGTTAGCAGCAAACCTGGATTCAATATGACAACCAACCAAGCTTCATGTGTCATCAAGTATGGACATT
 GAATTGTACACCATTAGTAGCCAACTAGATTCAATACGACTCGCAACCACACCATTAAAGTGTCTCATTAATATGGAGATG
 850 860 870 880 890 900 910 920

990 1000 1010 1020 1030 1040 1050 1060
 TAAGAGTGAATCAGACCTCAACTGGAATACAACCAAGCAAGAGCATTTCCTGATAACCTGCTCCATCCTGGCCATTAC
 CTCACGTGTCAGAGGACTTCACCTGGAAAAACCCCCAGAAGACCCCTCTGATAGCAAGAACACACTTGTGCTTTGGGGC
 930 940 950 960 970 980 990 1000

1070 1080 1090 1100 1110 1120 1130 1140
 CTTAATCTCAGTAAATGGAATTGTGATATGCTGCCTGACCTACTGCTTGCCTTCAAGATGCAGAGAGAGAAGGAGGAAT
 AGGATTGGCGAGTAATAACAGTCGTCATCGTTGTCAATCAAATGCTCTGTAAGCACAGAACAGCTGTTCAAGAAGA
 1010 1020 1030 1040 1050 1060 1070 1080

1150 1160 1170 1180 1190 1200 1210 1220 1230
 GAGAGATTGAGAAGGGAAAGTGTACGCCCTGTATAACAGTGTCCGCAGAACAGCAAGGGCTGAAAAGATCTGAAGGTAGCCTC
 AATGAGGCAAGCAGAGAAACAAACAACAGCCTTACCTTCGGGCCTGAAGAACAGCATTAGCTGAACAGACCGTCTTCCTTTAGT
 1090 1100 1110 1120 1130 1140 1150 1160 1170

1240 1250 1260 1270 1280 1290 1300 1310
 CGTCATCTCTGGGATACATGGATCGTGGGATCATGAGGCATTCTCCCTAACAAATTAAAGCTGTTTACCCACTAC
 TCTTCTCTGTCCATGTGGGATACATGGTATTATGTGGCTCATGAGGTACAATCTTCTTCAGCACCGTGCTAGCTGATCTT
 1180 1190 1200 1210 1220 1230 1240 1250

1320 1330 1340 1350 1360 1370 1380 1390
 CTCACCTCTTAAAAACCTCTTCAGATTAAGCTGAACAGTTACAAGATGGCTGGCATCCCTCTCCTTCTCCCCATATGCA
 TCGGACAACCTGACACAAGATAGAGTTAAGCTGGAAAGAGAAAGCCTTGAATGAGGATTCTTCCATCAGGAAGCTACGGGC
 1260 1270 1280 1290 1300 1310 1320 1330

1400 1410 1420 1430 1440 1450 1460 1470
 ATTTGCTTAATGTAACCTCTTCTTGCATGTTCCATTCTGCCATCTGAATTGCTTGTCAAGCCAATTCAATTCTATT
 AAGTTTGCTGGCCCTTGATTGCTTGATGACTGAAGTGGAAAGGCTGAGCCCAGTGTGGGTGGTGCTAGCCCTGGGAGGGG
 1340 1350 1360 1370 1380 1390 1400 1410

1480 1490
 AAACACTAATTGAG
 CAGGTGACCCCTGGGTGGTATAAGAAAAAGAGCTGTCACTAAAGGAGAGGTGCCTAGTCTTACTGCAACTTGATATGTCATG
 1420 1430 1440 1450 1460 1470 1480 1490

1500 1510 1520 1530 1540 1550 1560 1570 1580
 TTGGTTGGTGTCTGTGGGAGGCCTGCCCTTTCTGAAGAGAACGGTGGGAGAGTGGATGGGATGGGGGGAGAGGAAAAGT
 1590 1600 1610 1620 1630 1640 1650 1660
 GGGGAGAGGGCCTGGGAGGGAGAGGGAGGGAGGGACGGGGTGGGGTGGGAAAAGTATGGTGGGATGTAACACGGATA

Figure 8A: CD40 nucleotide sequence comparison between human, murine and cattle sequences.

Contig	AGDVTCGGATGAGAGCCCTGGTGGTATCCCCGTATGATGGGVATCCTGTTGCCATCCTCTTGGTG
Human CD40	AGGATCGGCTGAGAGCCCTGGTGGTATCCCCATCATCTCGGGATCCTGTTGCCATCCTCTTGGTG
Bovine CD40	AGAGTCGGATGAGGACCCCTGGTGGTATCCCCGTACGATGGGACTCTGTTGCTGTCCTGTTGGTA
Mouse CD40	AGTCCCGGATGGAGCCCTGCTGGTCACTCCTGTCGTATGGGATCCTCATCACCAATTCCGGGGTG
	690 700 710 720 730 740
Contig	TTTGTCTDTATC AAAAGGTGGCAAGAAGCCAACVGATAA NNNGCCCTVACCCCTANGCTNNANG
Human CD40	CTGGTCTTATCA AAAAGGTGGCAAGAAGCCAACCAATAA -----GCC CCCC CA CCCC A-----A
Bovine CD40	TCTGCCTGTATCAGGAACATAACCAAGAAGC-GGCAGCTAA-----GCC CTG ACCC T ATGGCTGAAAG
Mouse CD40	TTTCTCTATATCA AAAAGGTGGTCAAGAAACCAAGGATAATGAGATGTTACCCCTGCGCTCGAC G
	750 760 770 780 790 800 810
Contig	GCAGGATCCCCAGGAGATGAN T NCNGAVGAT TTT CCCGGCCCCAACACCGCTGCTCCAGTGCAGG
Human CD40	GCAGGAAC CCC AGGAGATCA TTT CCGAC GAT CT C CTGGCT CC AA ACT CGT G CTCCAGTGCAGG
Bovine CD40	GCAGGAT CCC TGGAGACGATTGAT CCGAGG AT TTT CCCGGCCCCAC-CCG CT CT CCGGT <u>GCAAG</u>
Mouse CD40	GCAAGATCCCCAGGAGATG-----GAAGATTAT CCCGT CATAACACCGCTGCTCCAGTGCAGG
	820 830 840 850 860 870 880
Contig	AGACHTTACACGGGTGT CAGCCGGT CACCCAGGAGGATGGCAAAGAGAG GTCG CATCTCAGT <u>TCAG</u> TGCAGGAG
Human CD40	AGACTTTACATGGATGCCAACGGT CACCC AGGAGGATGGCAAAGAGAG GTCG CATCTCAGT <u>TCAG</u> TGCAGGAG
Bovine CD40	AGACCTTATGCTGGTGT CAGCCGGT CGCCCAGGAGGACGGCAAAG
Mouse CD40	AGACACTGCACGGGTGT CAGC CTGT CACAC AGGAGGATGGTAAAGAGAG GTCG CATCTCAGT <u>TCAG</u> TGCAGGAG
	890 900 910 920 930 940 950
Contig	CGGCAGGTGACAGACAGC ATAGC CTTGAGGCC CTGGT CTGMACC CTGG A ACY G CTT YR RR GYGATG
Human CD40	-----AGACAG-----TGAGGC-----TG CACCC -----ACC-----CAGGAGTG-TG
Mouse CD40	CGGCAGGTGACAGACAGC ATAGC CTTGAGGCC CTGGT CTGAACC CTGG A ACT G CTT GGAGGCGATG
	960 970 980 990 1000 1010 1020
Contig# 1	GCYRCTTGCTGACCTTGAAG TTTGAG RTGRGCCAARACAGAGCCCAGTGCAGY TRRCY CTCATGCCT
Human CD40	GCCAC-----GTGGGC-----AAACAG-----GCAGTTGGCC-----
Mouse CD40	GCTGCTTGCTGACCTTGAAG TTTGAG ATGAGCCAAGACAGAGCCCAGTGCAGCTAACTCTCATGCCT

Contig	10	20	30	40	50	60
bovine CD40 protein
human CD40 protein	MVRLPLQCLFWGFFLTAVHSEPATACGEKQYPVNSLC	CDLCPPGQKL	VNDCTEVSKTECQ			
murine CD40 protein	MVRLPLQCVLWGCLL	TAVHPEPPTACREKQYL	INSQC	CSLCQPGQKL	VSDCTEFTETECL	
Contig	70	80	90	100	110	120
bovine CD40 protein
human CD40 protein	SCKGGEFLSTWNRE	KYCHEHRYCNPNLGLR	IQSEGTLNTDT	ICVCVEGQHCT	SHTCESCT	
murine CD40 protein	PCGESEFLDTWNRE	THCHQHKYCDPNLGLR	VQQKGTSETDT	CTCEEGWHCT	SEACESCV	
Contig	130	140	150	160	170	180
bovine CD40 protein
human CD40 protein	PHSLCLPGFGVKQIAT	GLLD	TVCEPCPLGFF	SNVSSAFEK	CHRWTSCERKGL	VEQHVG
murine CD40 protein	LHRSCSPGFGVKQIAT	GVD	TCEPC	PVGFF	CHPWTS	ETKDLVQQAGTN
Contig	190	200	210	220	230	240
bovine CD40 protein
human CD40 protein	KTDVVCFGQSRM	RTL	VV	IPV	TMG	VLFAV
murine CD40 protein	KTDVVCGPQDRL	RAL	VVI	IIFG	GILFA	LLVFIKKVAKK
Contig	250	260	270	280		
bovine CD40 protein	WLKGRIPWRR	LL	IRRIFPA	--	PTRLSGARD	MLVSAGRPGGRQ
human CD40 protein	NFPDDLPGSNTA	APVQETL	HGCQPV	TQEDGKESRIS	RVQER	
murine CD40 protein	EDYPGHN	TAAPVQETL	HGCQPV	TQEDGKESRIS	VQERTDSIALRPLV	

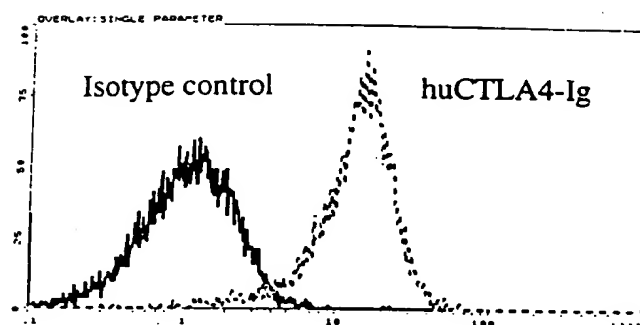
Figure 8B: Amino acid comparison between human, murine and cattle CD40 sequences.

A

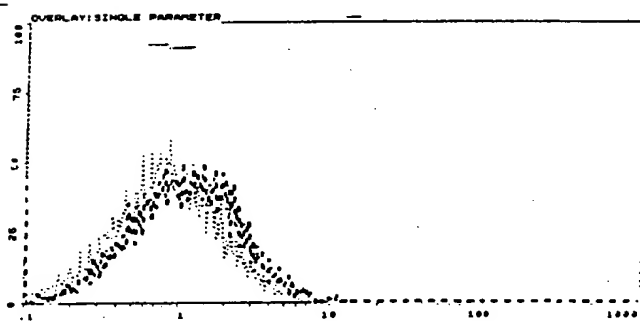
Non-transfected control cells



Transfected cells



Non-transfected control cells



Transfected cells

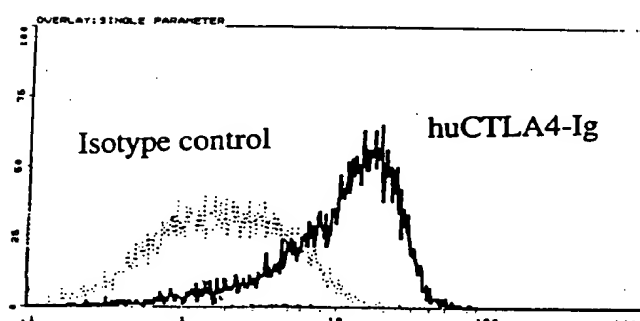
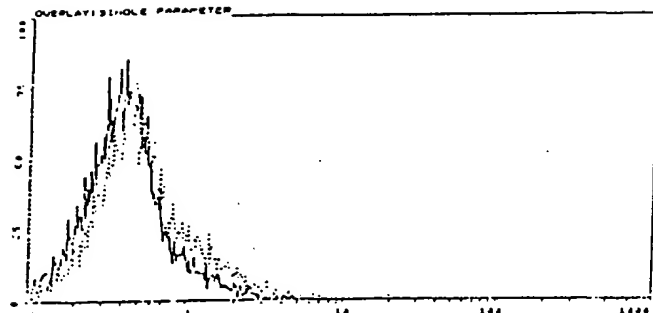
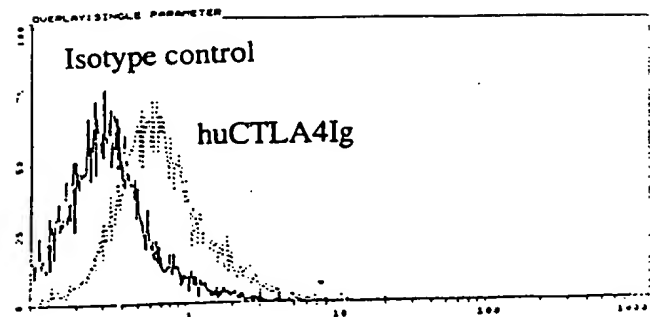


Figure 9: M1-poB7-2 (A) and P815-poB7-2 (B) clones generated by calcium phosphate transfection followed by dynabead selection and cloning by limiting dilution. Expression of B7-2 on the surface of transfected or control cells as determined by fluorescence activated cell sorting. 2.5×10^5 cells were stained with Mab to B7-2 (huCTLA4Ig) or isotype control (huIg) at 1 g/ml. After washing, cells were incubated with goat anti-mouse Ig-FITC conjugate, fixed with 1% paraformaldehyde and analysed on a Coulter counter.

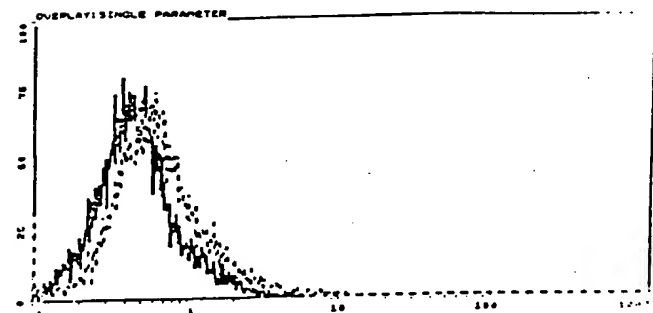
Non-transfected control cells



Transfected cells



Non-transfected control cells



Transfected cells

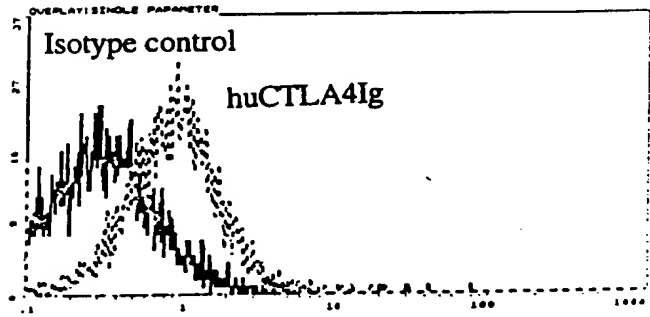


Figure 10: Transient transfections of M1 (A) and P815 (B) cells with CD86(i) by calcium phosphate precipitation. Surface expression of B7-2 on transfected or control cells was determined by fluorescence activated cell sorting. 48 hours after transfection, 2.5×10^5 cells were stained with Mab to B7-2 (huCTLA4Ig) or isotype control (huIg) at 1 g/ml. After washing, cells were incubated with goat anti-mouse Ig-FITC conjugate, fixed with 1% paraformaldehyde and analysed on a Coulter counter..

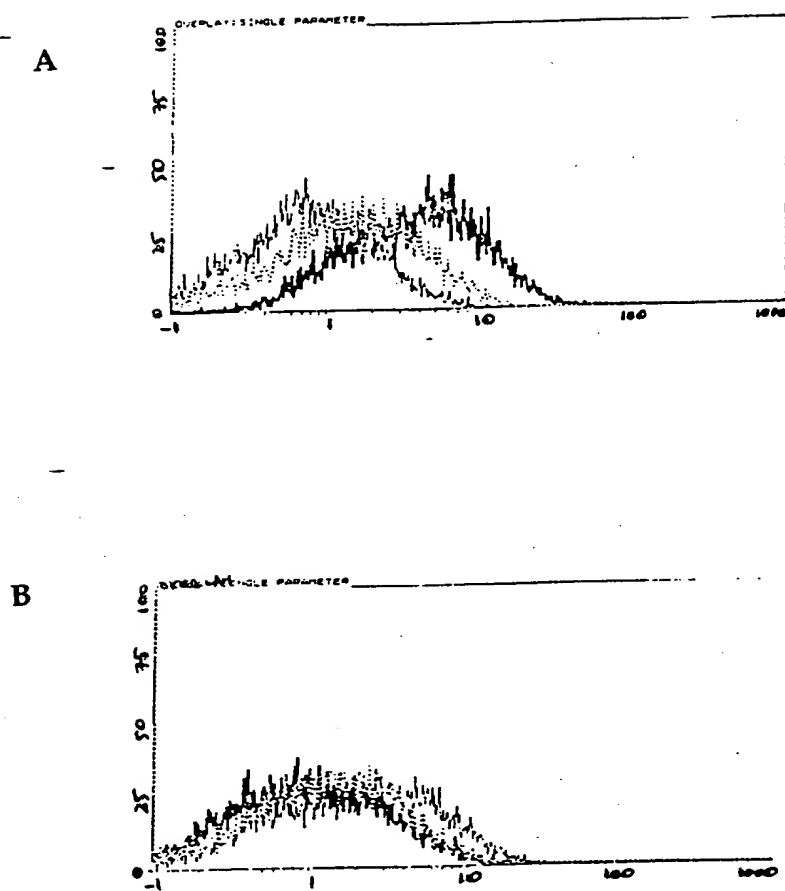
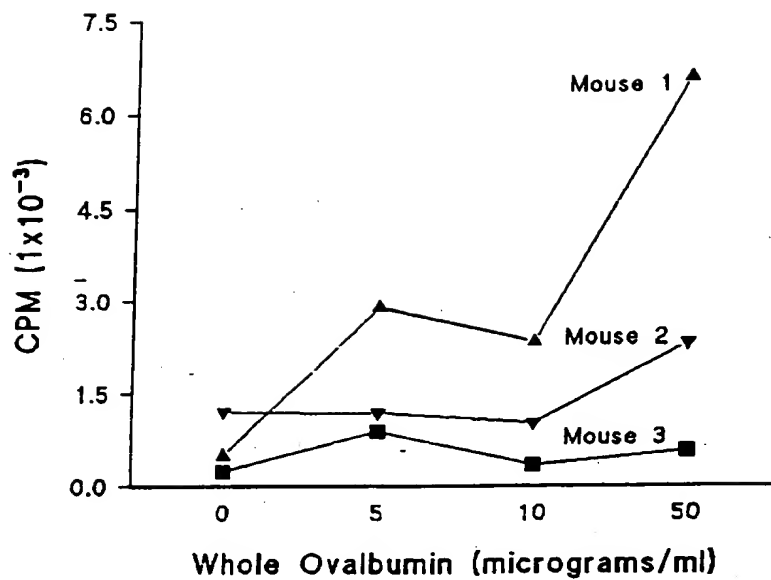


Figure 11: Flow cytometric analysis of porcine B7-2 transfected P815 cells following staining with porcine B7-2-specific sera or ovalbumin peptide control sera. 2.5×10^5 P815 cells were stained with 1/100 of each sera from B7-2 peptide (A) or ova control peptide (B) immunised mice. After washing, cells were incubated with goat anti-mouse IgG (H & L)-HRP and subsequently, Streptavidin-FITC. Cells were fixed with 1% paraformaldehyde and analysed on a Coulter counter.

1 MGLSNILFVM VLLLSGAASL KSQAYFNETG ELPCHFTNSQ
 9 | 8 | 2
 41 NLSLDELVIF WQDQDNLVLY ELYRGQEKPH NVNSKYMGRT
 10 |
 81 SFDQATWTLR LHNVQIKDKG SYQCFIHHKG PHGLVPIHQM
 1 | 5 | 3 |
 121 SSDLSLLANF SQPEINLLTN HTENSVINLT CSSTQGYPEP
 1 | 6 |
 161 QRMYMILLNTK NSTTEHDADM KKSQNNITEL YNVSIRVSLP
 1 |
 201 IPPETNVSIV CVLQLEPSKT LLFSLPCNID AKPPVQPPVP
 241 DHILWIAALL VTVVVVCGMV SFVTLRKRKK KQPGPSNECG
 281 ETIKMNRKAS EQTKNRAEVH ERSDDAQCDV NILKTASDDN
 321 STTDF•LKSK L

Figure 12: Positions of the nine B7-2 peptides with respect to the predicted amino acid sequence of porcine B7-2

A



B

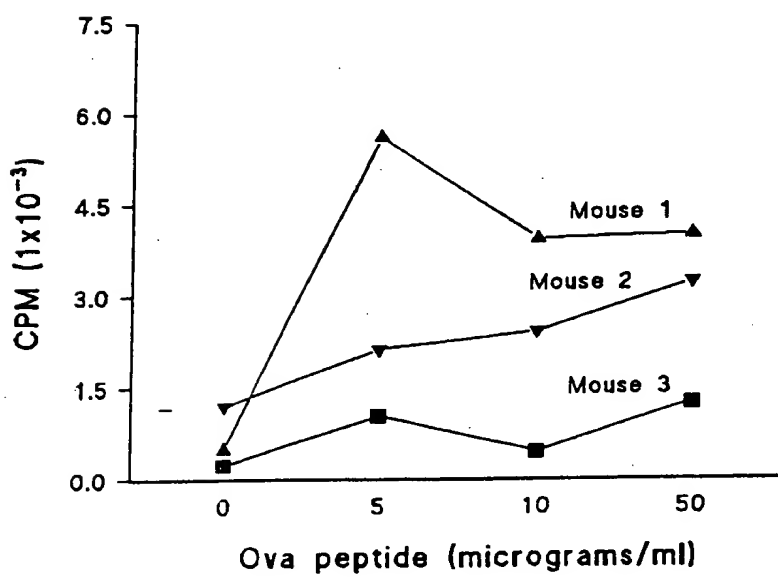
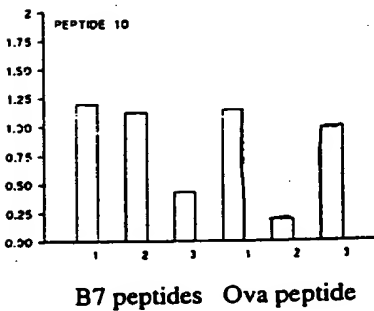
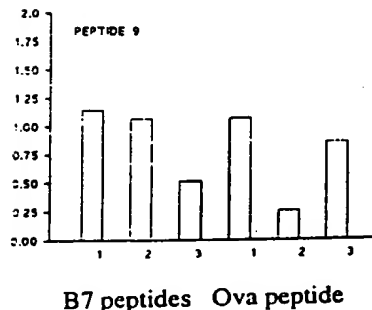
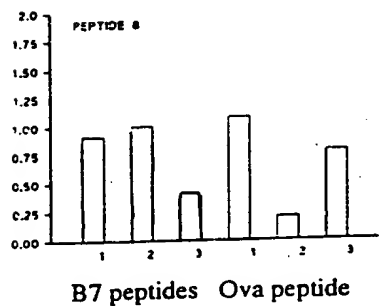
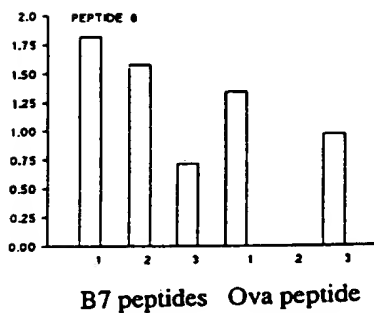
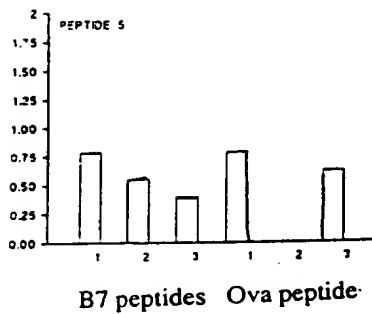
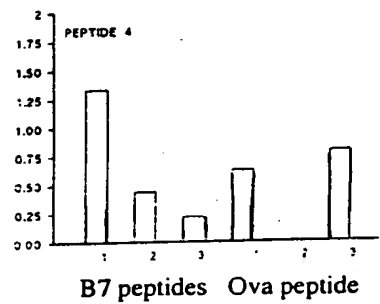
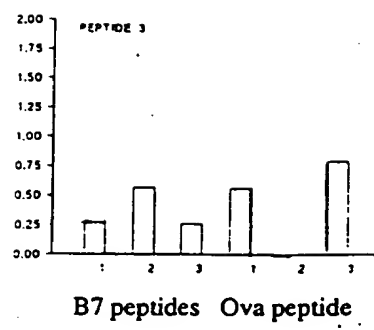
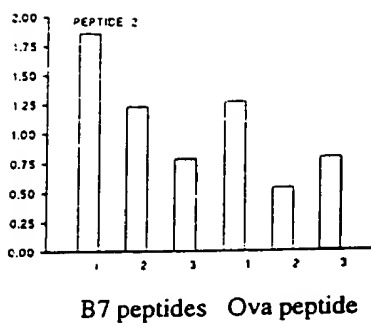
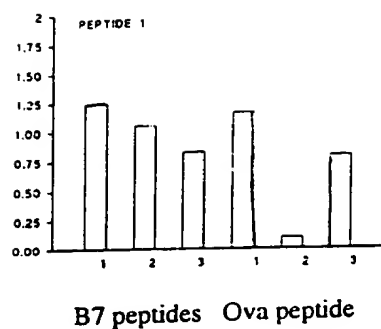


Figure 13: Comparison of *in vitro* T cell proliferation response to whole ovalbumin (A) or Ova₁₂₃₋₁₃₉ peptide (B). 2.5 x 10⁵ T cells and 2.5 x 10⁵ APC were plated per well with the indicated concentrations of whole ovalbumin or ova peptide. Cells were cultured for 72 hours in a total volume of 200 l 10% RPMI. T cell proliferation was assayed by the incorporation of ³H-thymidine.

Optical Density (450nm)



1:300 dilution of sera from immunised mice

Figure 14: Differential binding of B7-2 specific peptide sera or ova control sera as determined by Peptide ELISA. 96 well plates were pre-coated with the nine individual B7-2 specific peptides (P1-6 ; P8-10). Sera harvested from 3 individual B7-2 peptide (Bars 1-3), or 3 individual Ova control peptide (Bars 4-6) immunised mice were then screened for binding. Sera were detected by subsequent incubations with goat anti-mouse IgG-Biotin, Streptavidin-HRP and then developed with TMB. Plates were read at 450nm. Values have been adjusted for binding to no-peptide control plate and represent means for duplicate wells.

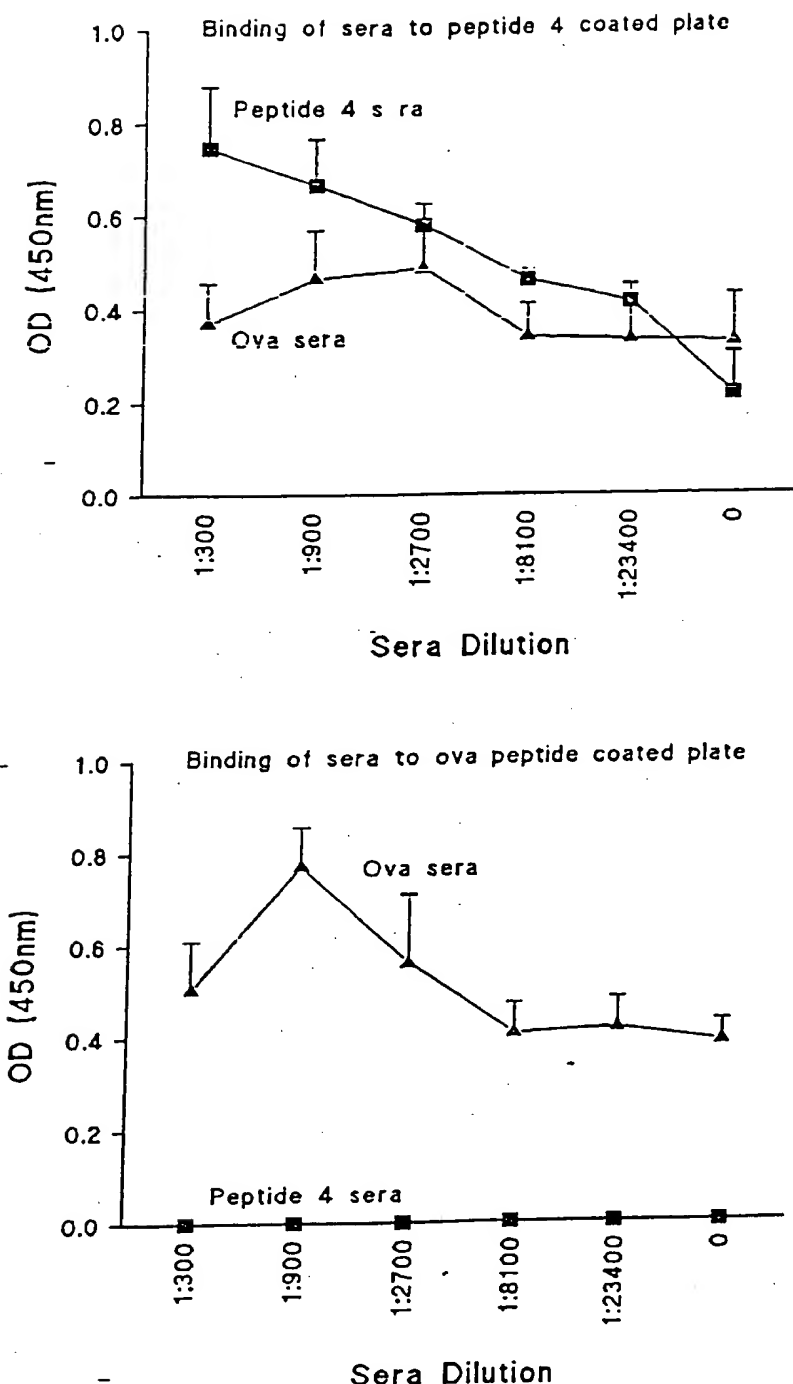


Figure 15: Differential binding of B7-2 specific sera or ova control sera as determined by Peptide ELISA. 96 well plates were pre-coated with either the B7-2 specific peptide by Pep4, Ova control peptide (OVA) or no peptide. Sera harvested from peptide 4, or Ova peptide immunised mice were then screened for binding. Sera were detected by subsequent incubations with goat anti-mouse IgG-Biotin, Streptavidin-HRP and then developed with TMB. Plates were read at 450nm.: Values represent means +/- SEM for 4 mice per group, in duplicate wells. Values have been adjusted for binding to no peptide control plate. Sera were measured over a range of dilutions.

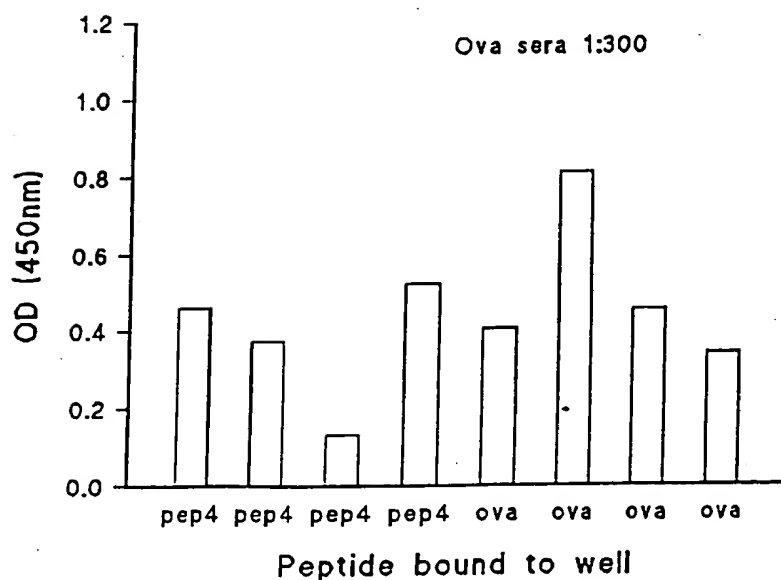
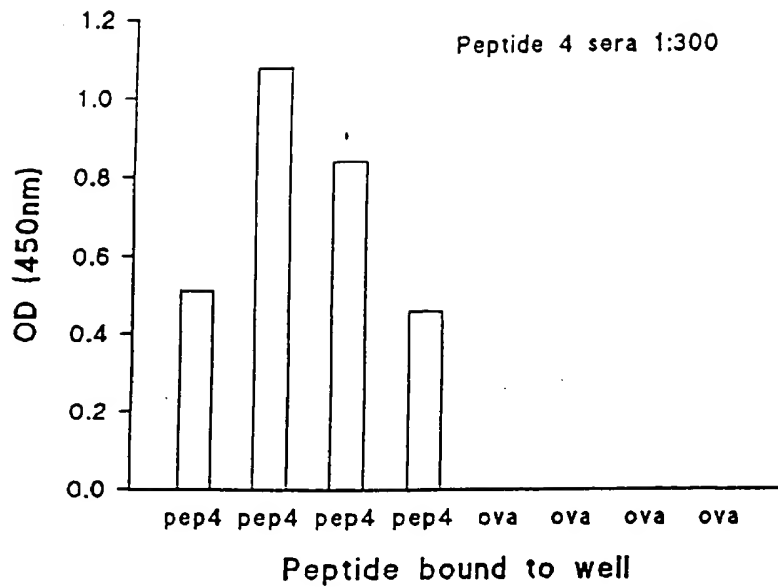


Figure 16 : Differential binding of B7-2 specific sera or ova control sera as determined by Peptide ELISA. 96 well plates were pre-coated with either the B7-2 specific peptide Pep4, Ova control peptide (OVA) or no peptide. Sera harvested from peptide 4, or Ova peptide immunised mice were then screened for binding. Sera were detected by subsequent incubations with goat anti-mouse IgG-Biotin, Streptavidin-HRP and then developed with TMB. Plates were read at 450nm. Values have been adjusted for binding to no-peptide control plates and represent means for duplicate wells for individual mice at 1:300 dilution of the sera.

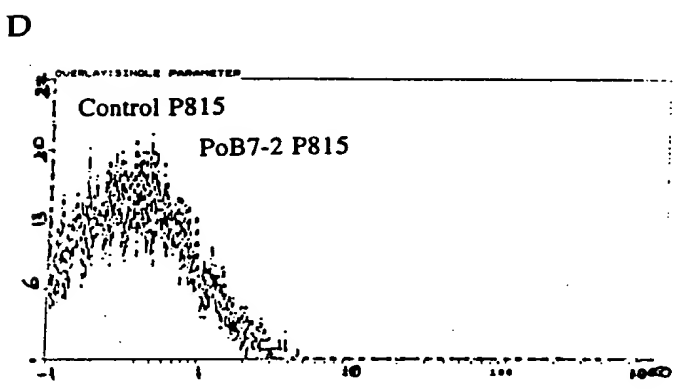
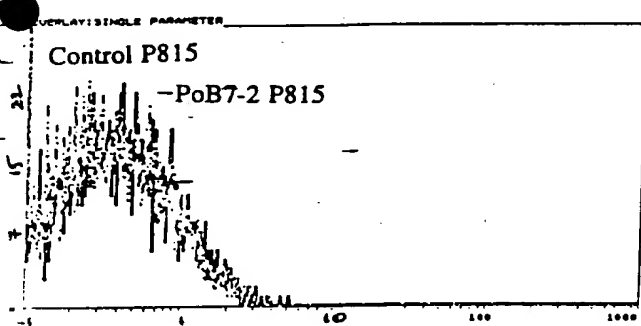
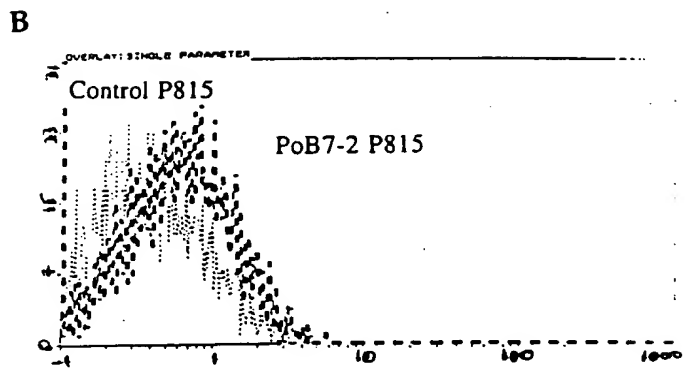
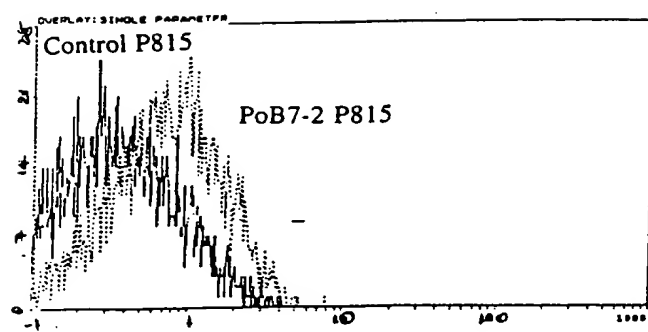


Figure 17: Flow cytometric analysis of porcine B7-2 transfected, or control untransfected P815 cells following staining with sera from peptide 4 or ovalbumin peptide control sera. 2.5×10^5 P815 cells were stained with 1 μ l of sera from 4 different mice immunised with either B7-2 peptide 4 (Figures A & B) or ova control peptide sera (Figures D & E). After washing, cells were incubated with goat anti-mouse IgG (H & L)-HRP and subsequently, Streptavidin-FITC. Cells were fixed with 1% paraformaldehyde and analysed on a Coulter counter.